

FISH & RICHARDSON P.C.



January 6, 1999

Attorney Docket No.: 08609/003004

Box Patent Application

Assistant Commissioner for Patents Washington, D.C. 20231

BOSTON

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TWIN CITIES

washington, DC Presented for filing is a new continuation patent application of:

Applicant:

LAWRENCE J. WANGH

Title:

PRENATAL SCREENING

The prior application is assigned of record to Brandeis University, a Massachusetts corporation, by virtue of an assignment submitted to the Patent and Trademark Office for recording on March 14, 1994 at 6931/126.

Enclosed are the following papers, including those required to receive a filing date under 37 CFR §1.53(b):

	rages
Specification	84
Claims	11
Abstract	1
Declaration	2
Drawing(s)	4

Enclosures:

- Copy of Power of Attorney (Revocation of Prior Powers), filed on parent application Serial No. 08/455,981, 2 pages.
- Copy of Small entity statement filed in parent application Serial No. 08/190,771, to which this application is entitled to priority under 35 U.S.C. §120. This application is entitled to small entity status.
- Rule 63 declaration, copy from a previous application under rule 63(d) for continuation or divisional only.
- Preliminary amendment, 11 pages.
- Postcard.

FISH & RICHARDSON P.C.

January 6, 1999 Page 2

This application is a continuation (and claims the benefit of priority under 35 USC 120) of U.S. application serial no. 09/050,380, filed March 30, 1998, which is a continuation of U.S. application serial no. 08/455,981 (Patent No. 5,773,217), filed May 31, 1995, which is a division of U.S. application serial no. 08/190,771 (Patent No. 5,651,992), filed February 1, 1994 which is a continuation-in-part of U.S. application serial no. 13,039 (Patent No. 5,480,772), filed February 3, 1993. The disclosure of the prior applications is considered part of (and is incorporated by reference in) the disclosure of this application.

Basic filing fee	380.00
Total claims in excess of 20 times \$9.00	450.00
Independent claims in excess of 3 times \$39.00	0.00
Fee for multiple dependent claims	0.00
Total filing fee:	830.00

A check for the filing fee is enclosed. Please apply any other required fees or any credits to deposit account 06-1050, referencing the attorney docket number shown above. If this application is found to be incomplete, or if a telephone conference would otherwise be helpful, please call the undersigned at 212/765-5070. Please acknowledge receipt of this application by returning the enclosed postcard.

Please send all correspondence to:

William J. Hone Fish & Richardson P.C. 45 Rockefeller Plaza New York, NY 10111

Respectfully submitted,

William J. Hone

Reg. No. 26,739

Enclosures

42538.N11

Applicant or Patentee: Lawrence J. Wangh Serial or Patent No.: 08/190,771

Filed or Issued: For:

February 1, 1994 PRENATAL SCREENING

VERIFIED S (37 CF	TATEMENT (DECLARATION) CLAIMING SMALL ENTITY STATUS FR 1.9(f) and 1.27(c)) - SMALL BUSINESS CONCERN
I hereby declare th	hat I am:
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a small business co 1.9(d), for purpose 35, United States including those of of this statement, average over the pr a full-time, part-t fiscal year, and (2 or indirectly, one	nat the above-identified small business concern qualifies as oncern as defined in 13 CFR 121.3-18, and reproduced in 37 CFR so of paying reduced fees under Section 41(a) and (b) of Title Code, in that the number of employees of the concernits affiliates, does not exceed 500 persons. For purposes (1) the number of employees of the business concern is the revious fiscal year of the concern of the persons employed or time or temporary basis during each of the pay periods of the concern are affiliates of each other when either, directly concern controls or has the power to control both.
remain with the sm	hat rights under contract or law have been conveyed to and mall business concern identified above with regard to the ention, by inventor(s) LAWRENCE J. WANGH described in
X a	the specification filed herewith pplication serial number 08/190,771, filed February 1, 1996 Patent No, issued
exclusive, each in invention is listed other than the inve 37 CFR 1.9(d) or h concern under 37 C *NOTE: Separate	d by the above-identified small business concern are not individual, concern or organization having rights to the below and no rights to the invention are held by any person, intor, who could not qualify as a small business concern under by any concern which would not qualify as a small business FR 1.9(d) or a nonprofit organization under 37 CFR 1.9(e) verified statements are required from each named person, at ion having rights to the invention averring to their status (37 CFR 1.27)
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I acknowledge the duty to file, in this application or patent, notification of any change in status resulting in loss of entitlement to small entity status prior to paying, or at the time of paying, the earliest of the issue fee or any maintenance fee due after the date on which status as a small entity is no longer appropriate. (37 CFR 1.28(b))
I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application, any patent issuing thereon, or any patent on which this verified statement is directed.
NAME OF PERSON SIGNING: JOEL M. COHEN, Director
NAME OF PERSON SIGNING: TITLE OF PERSON OTHER THAN OWNER: ADDRESS OF PERSON SIGNING: JOEL M. COHEN, Director Grant, Contract & Patent Admin.
signature Joel Cohen Date 2/28/94

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant: Lawrence J. Wangh Art Unit: Serial No.: Examiner:

Filed

Title : PRENATAL SCREENING

Box Non Fee Amendment

Assistant Commissioner for Patents Washington, DC 20231

PRELIMINARY AMENDMENT

Before calculating the filing fee, please amend the application as follows:

In the specification:

Delete the first sentence of the specification, and insert as the first sentence:

--This application is a continuation of U.S. application Serial No. 09/050,380, filed March 30, 1998; which is a continuation of U.S. application Serial No. 08/455,981, filed May 31, 1995, now U.S. Patent No. 5,773,217; which is a division of U.S. application Serial No. 08/190,771, filed February 1, 1994, now U.S. Patent No. 5,651,992; which is a continuation-in-

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part of U.S. application Serial No. 13,039, filed February 3, 1993, now U.S. Patent No. 5,480,772.--

In the claims:

Cancel claims 1-86 and substitute therefor the following claims:

87. In transplantation of a donor-cell nucleus other than a sperm or egg nucleus into a recipient egg for the purpose of creating a new organism capable of development, wherein said donor-cell nucleus, due to its gene structure and function, lacks the capacity to give rise to a complete, whole animal if directly transplanted into an activated recipient egg, the improvement comprising incubating said donor-cell nucleus in cytoplasms consisting of two types: first, in cytostatic factor-containing cytoplasm that is arrested in metaphase selected from the group consisting of meiotic metaphase II and mitotic metaphase; and subsequently in activated-egg cytoplasm.

- 88. The method of claim 87 further comprising step of removing or destroying said recipient egg's own nucleus.
- 89. The method of claim 88 wherein the step of removing or destroying the recipient egg's own nucleus precedes transplantation.
- 90. The method of claim 87 wherein the cytoplasm that is arrested in metaphase is a first extract from at least one egg.

- 91. The method of claim 90 wherein said first extract has been frozen.
- 92. The method of claim 90 wherein transplantation is by microinjection after the step of incubating with said first extract.
- 93. The method of claim 90 wherein said first extract is enriched for at least one of the group consisting of mitosis-promoting factor and cytostatic factor.
- 94. The method of claim 93 wherein said first extract is supplemented with at least one chemical or factor that enhances, stabilizes or extends the life of at least one kinase activity of said extract.
- 95. The method according to claim 94 wherein said first extract includes an aqueous buffer.
- 96. The method of claim 87 wherein the activated-egg cytoplasm includes a second extract from at least one egg.
- 97. The method of claim 96 wherein said second extract has been frozen.
- 98. The method of claim 97 wherein the cytoplasm that is arrested in metaphase is a first extract from at least one egg, and said first extract has been frozen.
- 99. The method of claim 96 wherein transplantation is by microinjection after the step of incubating with said second extract.

- 100. The method of claim 99 wherein said second extract has at least 70% activating capacity as determined by DNA replication.
- 101. The method of claim 100 wherein said second extract contains at least one inhibitor of an activating extract-promoted event or process.
- 102. The method of claim 87 wherein the activated-egg cytoplasm includes the recipient egg's cytoplasm.
- 103. The method of claim 102 wherein the recipient egg is treated so as to prevent activation of its cytoplasm prior to transplantation.
- 104. The method of claim 87 wherein the cytoplasm that is arrested in metaphase and the activated-egg cytoplasm are from different individuals.
- 105. The method of claim 104 wherein the different individuals are from different species.
- 106. The method of claim 87 wherein said donor-cell nucleus is a G2 nucleus.
- 107. The method of claim 87 wherein said cytoplasm that is arrested in metaphase is supplemented with at least one chemical or factor that enhances, stabilizes or extends the life of at least one kinase activity.
- 108. The method of claim 87 wherein the cytoplasm is arrested in metaphase and the activated-egg cytoplasm are cytoplasms of different eggs.

- 109. The method of claim 87 wherein said cytoplasm that is arrested in metaphase is supplemented with an inhibitor of kinase activity.
- 110. The method of claim 109 wherein said inhibitor is 6-dimethylamino purine (DMAP).
- 111. The method of claim 87 wherein, prior to contacting the donor-cell nucleus with the cytoplasm that is arrested in metaphase, the cytoskeleton surrounding the donor-cell nucleus is structurally altered.
- 112. The method of claim 111 wherein said cytoskeleton is structurally altered by at least one reagent selected from the group consisting of a thiol reducing agent, an ionic salt, an anionic chemical, a protease and a chelating agent.
- 113. The method of claim 111 further comprising step of removing or destroying said recipient egg's own nucleus.
- 114. The method of claim 113 wherein the step of removing or destroying the recipient egg's own nucleus precedes transplantation.
- 115. The method of claim 111 wherein the cytoplasm that is arrested in metaphase is a first extract from at least one egg.
- 116. The method of claim 115 wherein said first extract has been frozen.
- 117. The method of claim 115 wherein transplantation is by microinjection after the step of incubating with said first extract.

- 118. The method of claim 115 wherein said first extract is enriched for at least one of the group consisting of mitosis-promoting factor and cytostatic factor.
- 119. The method of claim 118 wherein said first extract is supplemented with at least one chemical or factor that enhances, stabilizes or extends the life of at least one kinase activity of said extract.
- 120. The method according to claim 119 wherein said first extract includes an aqueous buffer.
- 121. The method of claim 111 wherein the activated-egg cytoplasm includes a second extract from at least one egg.
- 122. The method of claim 121 wherein said second extract has been frozen.
- 123. The method of claim 122 wherein the cytoplasm that is arrested in metaphase is a first extract from at least one egg, and said first extract has been frozen.
- 124. The method of claim 121 wherein transplantation is by microinjection after the step of incubating with said second extract.
- 125. The method of claim 121 wherein said second extract has at least 70% activating capacity as determined by DNA replication.
- 126. The method of claim 125 wherein said second extract contains at least one inhibitor of an activating extract-promoted event or process.

- 127. The method of claim 111 wherein the activated-egg cytoplasm includes the recipient egg's cytoplasm.
- 128. The method of claim 127 wherein the recipient egg is treated so as to prevent activation of its cytoplasm prior to transplantation.
- 129. The method of claim 111 wherein the cytoplasm that is arrested in metaphase and the activated-egg cytoplasm are from different individuals.
- 130. The method of claim 129 wherein the different individuals are from different species.
- 131. The method of claim 111 wherein said donor-cell nucleus is a G2 nucleus.
- 132. The method of claim 111 wherein said cytoplasm that is arrested in metaphase is supplemented with at least one chemical or factor that enhances, stabilizes or extends the life of at least one kinase activity.
- 133. The method of claim 111 wherein the cytoplasm is arrested in metaphase and the activated-egg cytoplasm are cytoplasms of different eggs.
- 134. The method of claim 111 wherein said cytoplasm that is arrested in metaphase is supplemented with an inhibitor of kinase activity.
- 135. The method of claim 134 wherein said inhibitor is 6-dimethylamino purine (DMAP).

- 136. The method of claim 87 wherein said activated-egg cytoplasm is treated with a chemical that enhances its nuclear activation activity.
- 137. The method of claim 136 wherein said chemical is a protein synthesis inhibitor.
- 138. The method of claim 137 wherein the protein synthesis inhibitor is cycloheximide.
- 139. The method of claim 87 wherein said activated-egg cytoplasm is treated with a chemical that inhibits at least one event or process promoted by activated-egg cytoplasm.
- 140. The method of claim 139 wherein said chemical is a phosphodiesterase inhibitor.
- 141. The method of claim 87 wherein said donor-cell nucleus is the nucleus of a non-dividing cell.
- 142. A method for activating a nucleus of an animal cell comprising
- a) permeabilizing the plasma membrane of said cell, and
- b) incubating the nucleus of said cell in cytoplasms consisting of
- (i) cytostatic factor-containing egg cytoplasm that is arrested in metaphase selected from the group consisting of meiotic metaphase II and mitotic metaphase, and
 - (ii) activated egg-cytoplasm,

wherein incubation in said cytostatic factor-containing egg cytoplasm precedes incubation in said activated egg cytoplasm.

- 143. The method of claim 142 further comprising the step of transplanting said nucleus into a recipient egg.
- 144. The method of claim 143 wherein said animal cell is a somatic cell.
- 145. The method of claim 144 wherein said somatic cell is a human somatic cell.
- 146. The method of claim 144 further comprising the step of destroying said recipient egg's own nucleus.
- 147. The method of claim 142 wherein said animal cell is a sperm cell.
- 148. The method of claim 147 wherein said sperm cell is an human sperm cell.
- 149. The method of claim 147 wherein, prior to incubation in cytostatic factor containing cytoplasm, the cytoskeleton said nucleus is structurally altered.
- 150. The method of claim 149 wherein said cytoskeleton is structurally altered by at least one reagent selected from the group consisting of a thiol reducing agent, an ionic salt, an anionic chemical, a protease and a chelating agent.
- 151. The method of claim 142 wherein said activatedegg cytoplasm is treated with a chemical that enhances its nuclear activation activity.
- 152. The method of claim 151 herein said chemical is a protein synthesis inhibitor.

- 153. The method of claim 152 wherein the protein synthesis inhibitor is cycloheximide.
- 154. The method of claim 142 wherein said activatedegg cytoplasm is treated with a chemical that inhibits at least one event or process promoted by activated-egg cytoplasm.
- 155. The method of claim 154 wherein said chemical is a phosphodiesterase inhibitor.
- 156. The method of claim 142 wherein said donor-cell nucleus is the nucleus of a non-dividing cell.

REMARKS

Presented herewith are claims 87-156. Claims 87 and 142 are independent claims.

Claims 87-141 and 144-146 are limited to methods that exclude sperm cells, i.e., cloning.

Cloning claims have not previously been considered for patentability by the Office. Cloning claims were previously presented by Applicant in priority application Serial No. 08/455,891, but were subject to a restriction requirement in that case. Claims 142-143 and 151-156 are generic to somatic and sperm cells, and claims 147-150 are specific to sperm cells. None of this claims, however, is of the same scope as the claims of U.S. Patent 5,773,217, which claims are specific to activation

of sperm cells using CSF extract.

CONCLUSION

Applicants maintain that claims 87-156 patentably define the invention and request that the Examiner permit the pending claims to pass to allowance.

If the Examiner would like to discuss any of the issues raised in this Preliminary Amendment, the Applicant's undersigned representative can be reached at the telephone number listed below.

Enclosed is a check for the filing fee. Please charge any additional fees or make any credits to Deposit Account No. 06-1150.

Date: 1/6/99

Respectfully submitted,

Reg. No. 26,739

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42541.N11

APPLICATION

FOR

UNITED STATES LETTERS PATENT

TITLE:

PRENATAL SCREENING

APPLICANT:

LAWRENCE J. WANGH

"EXPRESS MAIL" Mailing Label Number <u>FE647285338US</u>

Date of Deposit <u>JANUARY</u> 6,1999

I hereby certify under 37 CFR 1.10 that this correspondence is being deposited with the United States Postal Service as "Express Mail Post Office To Addressee" with sufficient postage on the date indicated above and is addressed to the Assistant Commissioner for Patents, Washington, D.C. 20231.

PETER CONSOLI

DESCRIPTION

PRENATAL SCREENING

RELATED APPLICATION

This application is a continuation-in-part of U.S. Application Serial No. 08/013,039 entitled "Prenatal Screening" the whole of which is hereby incorporated by reference herein (including the drawings).

FIELD ON THE INVENTION

This invention concerns products, methods and apparatus for analysis of non-dividing mammalian cell nuclei, such as human fetal cell nuclei and mammalian sperm cell nuclei.

BACKGROUND OF THE INVENTION

Jackson, <u>Seminars in Perinatology</u> 15:49 (1991), describes various procedures for prenatal diagnosis, including procedures to diagnose diseases. These procedures involve analysis of the DNA present in early embryonic stages. Specifically, Jackson mentions the use of a polymerase chain reaction to amplify genes, and the possibility of testing occytes by polar body assay. According to Jackson:

20 "There are other conceivable embryo The approaches for prenatal diagnosis. trophectoderm may be obtained at later, multicellular embryonic stages when more cells might be obtained and induced to replicate in 25 tissue culture. . . . Another approach to early prenatal diagnosis is the recovery of fetal cells in the maternal circulation. This tantalizing possibility for a non-invasive method has been pursued for several years by groups in both the 30 United States and the United Kingdom. Both groups originally sought placental immunologic markers for identification and recovery of these cells. Several trophoblast antibodies were developed, some of which appeared to have relative

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specificity for the fetal cell. After sporadic reports of success, recent articles appear to indicate that these markers are insufficiently specific and actually are attached to maternal cells frequently enough to make this approach unworkable to date."

Bianchi et al., Proc. Natl. Acad. Sci. USA 87: describe isolating fetal nucleated 3279 (1990),erythrocytes in maternal blood using a monoclonal antibody 10 against the transferrin receptor. They state that they "were successful in detecting the Y chromosomal sequence in 75% of male-bearing pregnancies, demonstrating that it is possible to isolate fetal gene sequences from cells in maternal blood."

According to Roberts, Science 18:378 (1991), two procedures available for prenatal screening are chorionic villus sampling (CVS) and amniocentesis. Both these procedures have problems involving waiting time and risk of miscarriage, "estimated at 1% to 2% for CVS and 0.5% Supra. Roberts also points out a 20 for amniocentesis." procedure for analyzing nuclear DNA directly when cells are in interphase.

Lohka and Masui, Science 220:719 (1983), describe envelope nuclear formation of а inducing the 25 demembraned sperm of Xenopus laevis using a cell-free preparation from the cytoplasm of activated eggs of Rana pipiens.

Leno and Laskey, J. Cell Biology 112:557, (1991), performed experiments using erythrocytes from adult chickens. According to Leno:

"Coppock et al. (1989) [Supra] have reported that a pretreatment with trypsin was required for nuclear decondensation and DNA replication of extract. egg erythrocyte nuclei in Xenopus Trypsin pretreatment was not required for nuclear decondensation and DNA replication in our extracts."

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Gordon et al., Experimental Cell Research 157:409 (1985), describe "a system for the activation of human sperm using cell-free extracts from Xenopus laevis eggs." Similarly, an abstract, by Brown et al., J. Cell Biology 5 99:396a (1984), indicate that nuclear changes which occur during the early phases of fertilization can be stimulated by injecting isolated sperm nuclei into heterologous recipient eggs, or by incubating frog sperm nuclei in the presence of cell-free extracts from frog eggs. They state 10 that they found human sperm can be activated in vitro using Xenopus laevis frog egg extract to stimulate the early events of nuclear activation, including chromatin decondensation, nuclear enlargement and DNA synthesis.

SUMMARY OF THE INVENTION

The present invention concerns products and methods useful for causing non-dividing nuclei to activate nuclear of go through more steps one or (e.g., The featured products and methods are activation). particularly useful for activating human fetal cell nuclei 20 and mammalian sperm cell nuclei. "Activation" of a nondividing cell nucleus refers to one or more of the following activities: nuclear swelling, nucleic acid replication, and nuclear entrance into mitosis thereby (arrested metaphase chromosomes producing metaphase replicating chromosomes). 25 chromosomes or activation refers to activation wherein all of the activities occur.

Nucleic acids can be analyzed at the different stages of activation, brought about by the present such information useful invention, to obtain information about nucleic acid structure, sequences, number of copies of a nucleic acid sequence, and nuclear location of a nucleic acid. Analysis of nucleic acids can be carried out using techniques known in the art such as 35 in situ hybridization and karyotype analysis of metaphase chromosomes.

One particular advantage of the present invention is its use in prenatal diagnosis. Activation of fetal cell nuclei can be used to facilitate prenatal diagnosis of various human conditions. Nuclei from of all types of 5 human fetal cells including blood cells (such as red cells, white cells and other circulating cells of the fetus), as well as other types of fetal cells such as cells found in the amniotic fluid, or cells derived from syncytial or trophoblasts (such as placenta 10 trophoblasts), can be activated using the described products and methods. Preferably, the fetal cells to be activated are recovered from the blood or tissue of a pregnant woman rather than directly from the fetus or placenta, thereby decreasing the likelihood of discomfort 15 or harm to the fetus and/or mother by the diagnosis procedure.

"Activation activity" refers to the ability of an agent to bring about nuclear activation. Examples of agents which bring about nuclear activation include a non-20 activated cytostatic factor (CSF) extract and activating egg extract. Enhancement of activation activity refers to an increase in the activation activity which is brought about by an agent which causes nuclear activation. Examples of agents which enhance nuclear activation caused by an activating agent include CSF extract, purified components thereof, and proteases.

measured using be Activation activity can Such techniques include techniques known in the art. microscopic visualization of swollen nuclei, incorporation 30 of labelled nucleic acid precursors into newly synthesized nucleic acid, microscopic visualization of metaphase chromosomes, and in situ hybridization.

The featured methods include pretreating a nondividing human nucleus to enhance its ability to activate, bringing about complete or partial nuclear activation, and both bringing about and analyzing such nuclear activation on a microchamber microscope slide. Other useful methods

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disclosed include preparing products such as an activating egg extract, a CSF extract, and a modified CSF extract; the use of a protease pretreatment step in the activation of sperm; an activation assay; a retroviral integration 5 assay; and a procedure for cloning whole animals using activated nuclei.

The featured products including activating egg extract, CSF extract, kits containing these extracts, and a microchamber microscope slide useful in analyzing nuclear activation, are also claimed as part of the present invention.

The nucleus of a non-dividing fetal cell or a sperm cell is normally small, has condensed chromatin, and Specific nucleic acid does not replicate or divide. 15 sequences in the nucleus of these cells can be stained by fluorescent in situ hybridization methods if the target nucleic acid sequence is accessible to the probe. However, the small size of the nucleus can affect the accessibility of particular nucleic acid sequences and the information obtained from successful 20 amount of hybridization hybridization. Moreover, successfully obtained are limited in spacial resolution by the size of the nucleus. As a result, obtaining a reliable fluorescent signal can be difficult and the information obtained by fluorescent staining generally indicates only the presence or absence of accessible specific sequences, and possibly the number of such sequences per nucleus.

In the featured methods, the present invention 30 brings about one or more stages of nuclear activation: decondensation, nuclear swelling, chromatin replication, and formation of metaphase chromosomes. Genetic information can be obtained from each of these stages, which are characterized by changes in nuclear 35 structure and function. Useful information obtained from these stages of activation include facilitating the visualization of a particular chromosomal region using a

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probe by increasing the spacial resolution during swelling thereby increasing accessibility of the chromosomal region to the probe; detecting the number of a particular type of chromosome initially present by determining the increased number of the particular chromosome brought about by replication; and visualizing chromosomal morphology by staining metaphase chromosomes, including the presence of one or more sequences at specific locations within chromosomes.

Thus, in the first aspect, the invention features a method for causing a nucleus from a human fetal cell to activate. Activation is brought about by contacting a pretreated or, preferably, a further pretreated nucleus, with activating egg extract.

The present invention can be used to study fetal cell nuclei acid isolated by different procedures. For example, fetal cells can be obtained from circulating maternal blood, or by techniques such as amniocentesis or chorionic biopsy. Preferably, the fetal cell is obtained in a non-invasive manner (e.g., without disturbing the womb). Fetal cells such as erythrocytes and leukocytes cross the placenta and circulate transiently in maternal blood. Furthermore, trophoblasts which form the outermost placenta layer can pinch off and circulate in maternal blood. Trophoblasts typically end up trapped in the maternal lung capillary network.

Nuclear isolation and pretreatment is preferably carried out using mild conditions. Mild conditions are those which allows for nuclear isolation and pretreatment while causing the minimal amount of protein and nucleic acid damage. Using mild conditions helps maintain the integrity of the nucleic acid thereby decreasing artifacts during subsequent staining, and prevents premature protease activation thereby allowing subsequent protease treatment to occur under controlled conditions chosen to optimize such treatment.

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Preferably, nuclear isolation and pretreatment to release a nucleus from its surrounding cytoskeleton thereby forming a pretreated nucleus is carried out in two steps; (1) membrane permeabilization, and (2) separation 5 or alteration (e.g., denaturation or degradation) of cytoskeletal proteins and nuclear matrix proteins. steps may be carried out simultaneously or separately. Formation of a pretreated nucleus is preferably carried out under conditions minimizing nucleic acid damage and 10 damage to histones.

Membrane permeabilization, opens up the membrane treatment. nuclear facilitating subsequent thereby membrane used for Different techniques may be permeabilization including hypotonic shock, shearing and Preferably a non-ionic detergent is used to 15 detergent. permeabilize the plasma and nuclear membranes. More used as the non-ionic preferably, lysolecithin is detergent.

Different procedures can be use to separate, proteins degrade the cytoskeletal denature, and surrounding the nucleus and nuclear matrix proteins within the nucleus. These procedures include the use of a thiol reducing agent to denature nuclear protein, using selectively salt extraction to remove controlled 25 cytoskeletal and nuclear matrix proteins, and using controlled poly-anionic treatment to facilitate separation of negatively charged nucleic acid from the positively charged nuclear proteins. Separation conditions should be chosen to ensure a minimal amount of damage to nucleic non-cytoskeletal proteins. histones, and acids, Preferably, a protease is used under mild conditions to remove cytoskeletal proteins surrounding the nucleus. More preferably, trypsin is used as the protease. most preferred embodiment, pretreatment is achieved using 35 trypsin and lysolecithin.

Activating egg extracts are used to bring about nuclear activation. Activating egg extracts contain

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material, such as precursors, protein(s), nuclear envelope vesicles and mRNA, which support nuclear activation. egg can be chemically, physically, or electrically induced to produce material which brings about nuclear activation. Eggs can be induced using a calcium ionophore as described The induced egg continues in its cell cycle. appears that when an egg is at the point in the cell cycle just prior to the S-phase, the egg cytoplasm is most active in supporting activation. As the egg proceeds into 10 and past the S-phase, it appears to produce material inhibitory to activation.

Preferably activating egg extracts are prepared from Xenopus eggs. More preferably activating egg extract are prepared from eggs having an elevated DNA synthesis 15 activation activity. Activating egg extract prepared from Xenopus eggs induced for 10 minutes at 20°C contain approximately 59% of the optimal DNA synthesis activation activity of Xenopus eggs induced for 25 minutes at 20°C. At about 25-30 minutes at 20°C the Xenopus eggs are at 20 highest (optimal), or peak, DNA synthesis activation Xenopus eggs induced for 40 minutes at 20°C appear to have a lower DNA synthesis activation activity Thus, the present than the peak activation activity. invention discloses the use of induced eggs having an elevated DNA synthesis activation activity of 70% greater of the peak activation activity.

Activating extracts prepared from Xenopus eggs induced for 10 minutes or less at 20°C produce a lower However, rate of DNA replication in treated nuclei. activating extract prepared from Xenopus eggs induced for 10 minutes at 20°C appear to produce equivalent or greater nuclear swelling in treated nuclei than extracts prepared from Xenopus eggs induced for more than 10 minutes at 20°C.

preferably activating extract 35 More egg prepared from a number of eggs (e.g., 1,000 to 10,000), most or all of which have an elevated or peak DNA

synthesis activation activity. Obtaining a large number of eggs having a peak or elevated DNA synthesis activation activity is preferably achieved using hardened eggs which Hardened eggs are have been synchronously induced. 5 prepared by hardening the vitelline envelope surrounding the egg (described in detail below). Hardened eggs are less likely to spontaneously activate than soft nonhardened eggs.

Thus, by using hardened eggs a large number of 10 eggs can be collected and induced at the same time (synchronously induced). A given number of eggs synchronously induced should all be at or near the same point in their cell cycle at a given later time. A large number of eggs having an elevated DNA synthesis activation activity can 15 be obtained by inducing the eggs at one time, preparing the activating egg extract from all the eggs at Preferably the activating egg a second later time. extract is stored frozen. Freezing the extract allows a large amount of extract to be prepared at one time and 20 used at different later times.

Various supplements to activating egg extract have been found to increase the activation activity of the activating egg extract. These supplements include cell cycle regulatory proteins, cell cycle inhibitors, cAMP (preferably, between 0.1 and 1.0 mM, most preferably at 0.3 mM), and phosphodiesterase inhibitors (preferably caffeine, more preferably caffeine at a concentration between 0.1 and 10.0 mM, most preferably caffeine at a concentration of 1 mM).

In another preferred embodiment activation occurs under nuclear non-duplication conditions wherein the metaphase DNA, forms nucleus swells, replicates chromosomes and prepares to divide (i.e., enters mitosis), but segregation of sister chromatids is prevented by 35 inhibiting spindle formation. The inhibition of spindle formation prevents the division of the cell nucleus and the resulting separation of metaphase chromosomes.

Thus, under non-duplication conditions metaphase chromosomes are detectable for a longer time period and are provided in a "spread pattern." A "spread pattern" refers to the orientation of different chromosomes with Drugs such as nocodazole, 5 respect to each other. colchine, or colcemid can be used to inhibit spindle formation. Preferably nuclear non-duplication conditions is achieved by adding nocodazole to the activating egg More preferably, nocodazole is in an amount extract. 10 which will not inhibit DNA replication (e.g., less than 5 $\mu g/ml)$.

In other preferred embodiments, prior to being treated with the activating egg extract, the pretreated nuclei are further pretreated by contact with a CSF 15 extract, or a purified component of the CSF extract including a purified kinase or a purified phosphatase. By "purified" is meant the component is more concentrated (e.g., has a higher specific acitivity) than when present The desired purified kinase or in a CSF extract. 20 phosphatase can be obtained by purifying the enzymes from CSF fractions and assaying for activation activity. Further pretreatment with CSF is preferably carried out under conditions not resulting in nucleus activation. non-controlled occurring under Premature activation 25 conditions decreases the ability of CSF extracts to enhance activation because activation is occurring in CSF non-optimized conditions. under disadvantage of premature activation is that it produces a pool of nuclei activated at different times, which is more difficult to examine than nuclei activated at the same time.

CSF extracts can be used to increase nuclear activation upon subsequent contact with an activating agent. CSF extracts can be prepared from non-induced eggs (i.e., eggs arrested in meiotic metaphase II or activated eggs that have been arrested in mitotic metaphase). These extracts contain factors which aid in nuclear activation,

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such as CSF and mitosis promoting factor (MPF). MPF may bring about activation and visualization chromosome by stimulating chromosome condensation and inhibiting spindle assembly.

A preferred source of CSF extracts is Xenopus Isolation of CSF extract from Xenopus eggs is which do using "hardened eggs" facilitated Preferably, the CSF extract is spontaneously induce. stored frozen. Freezing the extract allows a large amount 10 of extract to be prepared at one time and used at different later times.

CSF extract is preferably supplemented with reagents such as β -glycerol phosphate, creatine phosphate, phosphocreatine kinase, and Ca2+ in amounts which improves 15 activation of nuclei in activating extract, without causing the start of the cell cycle prior to contact with Preferably, the CSF extract activation egg extract. contains Ca2+ in an amount which leads to an increase in the level of histone kinase or MPF activity without 20 initiating the cell cycle. The use of Ca2+ to supplement CSF extract is particular advantageous if the CSF extract The Ca²⁺ may be added before is frozen before use. freezing or after thawing.

Ca2+ is a cofactor for calmodulin activated protein kinases and may increase CSF activity by 25 increasing the level of phosphorylated topoisomerase II Topoisomerase II is a scaffold protein which activity. chromosome decondensation and condensation in possibly by anchoring chromatin loop domains. Wood and Ca²⁺ also 30 Earnshaw, <u>J. Cell Biology</u> <u>111</u>:2839 (1990). appears to increase the histone kinase level, which we have used as one measure of MPF activity.

As would be appreciated by one skilled in the art, the optimal amount of Ca2+ added to a CSF extract varies depending upon the presence of a Ca2+ chelator. Ca2+ concentration is preferably equal to or greater than 100 μM ; more preferably the Ca^{2+} concentration is between 100 μ M and 400 μ M. These preferred concentrations were determined using a CSF extract supplemented with 1 mM ethylene glycol-bis(β -aminoethyl ether)N,N,N'N'-tetraacetic acid (EGTA).

In another preferred embodiment, nuclei are activated under non-synthesis conditions which inhibit nucleic acid synthesis. As a result, the nucleus swells with or without formation of a nuclear envelope but does not replicate DNA or enter mitosis. The resulting 10 increased spacial resolution brought about by nuclear swelling facilitates the use of nucleic acid probes by making regions of nucleic acid more accessible. synthesis conditions, which nevertheless permit nuclear swelling may be achieved by the addition of reagents such as aphidicolin (e.g., 50-100 μ g/ml), 6-dimethylaminopurine 15 (e.q., at 5 mM), leupeptin (e.q., at dideoxycytidine triphosphate (e.q., 0.1 (Mm or dideoxythymidine triphosphate (e.q., 0.1 mM) activated egg extract, or to CSF extract which is then 20 contacted with an activated egg extract.

In another aspect, a non-dividing human nucleus is further pretreated for subsequent activation by contact with a purified protein kinase or a purified phosphatase which is present in a CSF extract. The purified protein kinase or purified phosphatase is in a purer form (e.g., more concentrated or more active) than that found in a CSF extract.

In a third aspect, the invention features a method for activating a non-dividing human nucleus by further pretreating the non-dividing human nucleus in CSF extract, prepared from hardened eggs, and then contacting these further pretreated nuclei with an activating egg extract prepared from synchronously induced hardened eggs.

In preferred embodiments the CSF extract is frozen before use and the activating egg extract is frozen before use.

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In other preferred embodiments, pretreated nuclei undergo further pretreatment in CSF extract involving a warm-then-cold incubation regime. Both the warm and cold steps increase activation of nuclei upon subsequent 5 contact with activating extract. Preferably, incubation is carried out at about 25°C for at least 30 minutes followed by incubation at about 4°C for at least 30 minutes. Less preferred, but still an effective incubation, is a warm regime at about 25°C for at least 30 minutes.

other preferred embodiments, thawed extract is supplemented with Ca2+ in an amount which does not start the cell cycle but improves nuclei activation. The Ca2+ should be in an amount which leads to an increase in the level of histone kinase or MPF without initiating the cell cycle.

In another aspect, methods are described for preparing an activating egg extract, from hardened eggs, which can cause non-dividing human nuclei cells to acti-The activating egg extracts are prepared from 20 vate. hardened eggs which have been synchronously induced such that the activating egg extract is prepared from eggs having an elevated DNA synthesis activation activity. Preferably synchronous induction is carried out using 25 eukaryotic cells, more preferably amphibian, yeast, human, echinoderm, mollusc, or fish, or chicken cells are used; eggs are used; even more preferably Xenopus preferably Xenopus eggs induced for more than 10 minutes are used; most preferably Xenopus eggs are induced for 25-30 minutes at 20°C.

In another aspect a method for inducing swelling in non-dividing nuclei is described. The method can be used to induce swelling in the absence of an activating extract and in the absence of DNA synthesis. 35 particular, CSF extract is supplemented with a protein kinase inhibitor and/or an aqueous solution.

a method for chromosome In another aspect, formation without DNA replication is described. method involves using a CSF extract supplemented with a cyclin such as cyclin-090 in an amount sufficient to 5 enhance nuclear envelope breakdown and nuclear chromosome The cyclin is thought to act by raising the formation. level of MPF activity in a CSF extract.

In another aspect, a method for activating a mammalian sperm cell nucleus is described. The method involves the steps of: (a) pretreating a sperm cell, using a membrane permeabilizer, a protease, and a thiol reducing agent to form a pretreated sperm cell; and (b) activating The method can be used to the pretreated sperm cell. Such studies can be study sperm from different mammals. 15 carried out, for example, to determination whether the sperm contains a particular gene or nucleic acid sequence which can be passed on during fertilization.

assays activation aspect, another In These assays can be used to measure different described. 20 stages of activation. A basic assay comprises isolating a nucleus, pretreating the nucleus, further pretreating the nucleus, contacting the further pretreated nucleus with an activating egg extract and measuring activation Measurement of activation activity can be activity. such techniques as standard usina 25 carried out newly labelled nucleotides into of incorporation synthesized nucleic acid and microscopic visualization of nuclear swelling and metaphase chromosome formation.

Other activation assays are performed by altering one or more of the steps of the basic assay. 30 instance, to assay for important factors in CSF extract, rather than using whole CSF extracts, fractions of the These fractions are obtained using extract can be used. standard purification techniques. Similarly, different 35 activating egg extract fractions can be studied.

In a preferred embodiment, a sperm activation assay, particularly useful to study human male fertility,

is described. Uses of the sperm activation assay include, determining the effect of handling sperm under different condition thereby obtaining optimal handling condition for subsequent in vitro fertilization, and testing the effect of possible male contraceptives on activation.

integration aspects viral other involving the use of a cell nucleus or a pseudonucleus are described. Viral integration into a cell nucleus can be assayed as follows: pretreating a cell nucleus to 10 separate the nucleus from its surrounding cytoskeleton; activating the pretreated nucleus and incubating with a viral integration complex containing viral nucleic acid; and measuring integration of viral nucleic acid into nucleic acid of the cell nucleus. The viral integration 15 complex containing viral nucleic acid can be added at times during nuclear pretreatment and different activation.

The viral integration assay using a pseudonucleus involves: a) constructing a pseudonucleus from a defined 20 DNA template; b) replicating the pseudonucleus; and c) incubating the pseudonucleus in the presence of a viral integration complex containing viral nucleic acid. This integration complex can be added at any time during pseudonucleus formation or replication. A pseudonucleus can be constructed, for example, by adding plasmid DNA to a CSF extract or an activating extract. The plasmid forms chromatin in the CSF extract but does not replicate until Ca²⁺ (1-4 mM) is added. Activation of the extract containing the pseudonucleus leads to nuclear envelope formation around the chromatin template and causes the chromatin to replicate.

for further a product In another aspect, The further nuclei is described. of pretreatment with extract **CSF** pretreatment product comprises 35 supplemented with Ca2+. Ca2+ is provided in an amount which increases nuclei activation upon subsequent contact with activating extract. Preferably, the CSF extract is also

phosphate, creatine with β -glycerol supplemented phosphate, and phosphocreatine kinase,

In preferred embodiments the CSF extract is frozen; the Ca2+ concentration is equal to or greater than 100 μ M; more preferably the Ca²⁺ concentration is between These preferred embodiments were 100 μ M and 400 μ M. determined using a CSF extract supplemented with 1 mM EGTA.

In another aspect, a product for causing nuclear The product contains CSF extract 10 swelling is described. supplemented with a protein kinase inhibitor and/or an aqueous solution.

product for In another aspect a chromosome formation without DNA replication is described. 15 The product is made up of a CSF extract supplemented with a cyclin such as cyclin-090 in an amount sufficient to bring about nuclear envelope breakdown and nuclear chromosome formation.

In another aspect, a product for causing a non-20 dividing nucleus to activate is described. The activating product comprises an activating egg extract prepared from an egg(s) having an elevated DNA synthesis activation activity.

In a preferred embodiment activating egg extract is prepared from Xenopus eggs synchronously induced for more than 10 minutes; preferably the Xenopus eggs are induced for 25 to 30 minutes at about 20°C.

In other preferred embodiments, the activating egg extract is modified by supplementation with cell cycle inhibitors, 30 regulatory proteins, cell cycle (preferably, between 0.1 and 1.0 mM, most preferably at 0.3 mM), and phosphodiesterase inhibitors (preferably caffeine, more preferably caffeine at a concentration between 0.1 and 10.0 mM, most preferably caffeine at a concentration of 1 mM).

In another aspect, a kit is disclosed for activating a non-dividing nucleus. The kit is comprised

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of frozen activating egg extract prepared from an egg having an elevated DNA synthesis activation activity and frozen CSF extract.

a preferred embodiment the CSF 5 contains Ca2+. In a most preferred embodiment the kit contains a microchamber microscope slide.

In another aspect, the invention features a microchamber microscope slide provided with an upper surface having a water-repellent material of a known 10 thickness defining a microchamber on the upper surface. The microchamber is shaped to enhance flushing of the microchamber, and connected by at least one channel to a well on the upper surface.

In preferred embodiments, the microchamber is teardrop-shaped or pear-shaped; preferably two wells are provided at opposite ends of the microchamber connected by two separate channels to the microchamber; and the microchamber has a defined volume preferably between 5 and 50 μ l, more preferably between 10 and 20 μ l when a 20 coverslip is placed over it. Fluid can be introduced into the microchamber by placing fluid in one well and allowing it to flow through the microchamber to the opposite well. The fluid is then removed from the opposite well. Removal may be achieved by pipetting away the fluid or by capillary action by placement of a filter paper within the well.

other preferred embodiments, the water-In repellent material is a tape or a coating on the upper surface of the slide, more preferably a TEFLON coating, or In most preferred a wax film (<u>e.g.</u>, a PARAFILM^{*}). embodiments, the upper slide surface is treated to enhance cell growth compared to an untreated slide, the slide is provided in a sterile condition, and/or the slide is coated with an antibody able to specifically bind to a 35 human fetal cell.

The advantages of the present invention include, but are not limited to, facilitating prenatal screening by

optimizing conditions for nuclear activation, which causes the nucleus of a fetal cell to either swell, replicate chromosomes. metaphase form nucleic acid. and/or Important information regarding nucleic acid sequences or 5 chromosome morphology can be readily obtained from these various stages of activation, for example by using DNA probes or visualizing the produced metaphase chromosomes. as fetal cells, such some erythrocytes, and leukocytes can be obtained from a 10 maternal source, an advantage of the invention is a noninvasive procedure to detect the presence of genetic defects in such cells.

Other features and advantages of the invention will be apparent from the following description of the preferred embodiments thereof, and from the claims.

Description of the Preferred Embodiments

The drawings will first briefly be described.

Drawings

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Fig. 1 shows the effect on DNA replication of 20 activated nuclei, of using frozen/thawed activating egg extracts supplemented with caffeine.

Fig. 2 shows the effect on DNA replication of activated nuclei, of using CSF extract supplemented with 6-dimethylamino-purine (DMAP).

25 Fig. 3 shows the effect of various warm-then-cold protocols on DNA replication in activated nuclei.

Fig. 4 is a top view of a microchamber microscope slide.

Methods And Products

Methods for activating nuclei include those described by Coppock et al., <u>Developmental Biology 131</u>:102 (1989); Wangh, <u>J. Cell Science 93</u>:1 (1989); Wood and Earnshaw, <u>J. Cell Biology 111</u>:2839 (1990); Leno and Laskey, <u>J. Cell Biology 112</u>:557 (1991); Young, <u>Biology of Reproduction 20</u>:1001 (1979); Philpott et al., <u>Cell 65</u>:569

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(1991); Shamu and Murray, J. Cell Biology 117:921 (1992); Adachi et al., Cell 64:137 (1991); Newport and Spann, Cell 48:219 (1987); and Henry Harris, in CELL FUSION 40-50 (Harvard University Press 1970).

DiBerardino et al., Proc. Natl. Acad. Sci. USA 83:8231 (1986), and Orr et al. Proc. Natl. Acad. Sci. USA (1986)describe nuclear transplantation experiments to activate Rana pipiens nuclei. DiBerardino was able to obtain tadpoles having a survival rate of up 10 to a month, by transplanting differentiated somatic cells into enucleated eggs.

The present invention discloses methods products useful in activating a non-dividing nucleus, and studying such activation. These methods and products are especially useful for analyzing a nucleus from nondividing fetal cells such as aminocytes, human keratinocytes, trophoblasts, erythrocytes and leukocytes. However, the methods and products are also useful for activating the nuclei from other types of non-dividing 20 human cells such as other types of non-dividing fetal cells and sperm, and non-dividing cells isolated from other mammals.

Preparing a nucleus for nuclear activation and bringing about nuclear activation is described in detail 25 below as four different phases: (1) preparation of nondividing human nuclei, (2) preparation of activating egg extracts from a source such as activated Xenopus eggs, (3) preparation of non-activated CSF extracts from a source such as non-activated Xenopus eggs, and (4) activation of non-dividing human nuclei.

Also described in detail below are modified CSF extracts which can bring about nuclear swelling in the absence of an activating egg extract; new procedures of pretreating a sperm cell to enhance its activation; a 35 microchamber microscope slide which facilitates bringing about nuclei activation and analysis of nucleic acids in such cells; a kit for bringing about nuclear activation;

an activation assay; and a procedure for cloning whole organisms from somatic cell nuclei.

The featured methods and products can be used to cause activation of a non-dividing human nucleus thereby 5 inducing swelling, and/or DNA replication and/or the formation of metaphase chromosomes. The procedures provided herein regarding nuclei activation are generally based upon existing procedures used in other systems. However, several improvements over the existing systems 10 are disclosed. Furthermore, existing procedures have not previously been used on human fetal cells nor was it known if they would produce useful results on such cells.

Examples are given to illustrate different aspects and embodiments of the present invention. 15 to be understood that various different modifications are possible and are contemplated within the true spirit and There is no intention, scope of the appended claims. therefore, of limitations to the exact process or disclosure herein presented.

the there is shown below particular, In activation of a human fetal red blood cell nucleus using frozen/thawed activating egg extract without further pretreatment, under non-dividing conditions. The treated nuclei swelled significantly, replicated DNA, and then 25 entered and arrested in the pre-mitotic state. nuclei are useful for prenatal diagnosis. Furthermore, the use of further pretreatment, should increase the rate and extent of nuclear swelling, decrease the time it takes for DNA synthesis to occur after activation, increase the 30 rate and extent of DNA synthesis, and improve the efficiency with which metaphase chromosomes are formed.

I. NUCLEAR ACTIVATION

(1) Preparation Of Nuclei

The present invention provides a method for 35 activation non-dividing mammalian cell nuclei, preferably

non-dividing human cell nuclei. Before being activated non-dividing human nuclei are isolated and pretreated. A preferred source of non-dividing human cells are fetal cells recovered from the blood of pregnant women such as leukocytes (such erythrocytes and 5 trophoblasts, granulocytes, neutrophils, basophiles and eosinophils). Isolating these cells does not require penetration of the womb. The present invention is also useful for analyzing other types of non-dividing human cell nuclei, including 10 non-dividing keratinocytes (e.g., those isolated from amniotic fluid), aminocytes, and sperm cells, or similar cells obtained from mammals other than humans.

Non-dividing fetal cells can be recovered from maternal blood supply using techniques such as antibody 15 staining followed by cell sorting. (For example, Bianchi entitled Non-Invasive Method For Isolation and PCT/US90/06623, Detection of Fetal DNA" incorporated by reference herein). Antibody cell sorting techniques separate fetal and maternal cells based on the presence of different antigens on fetal and maternal The antigen can be differentiated by suitable antibodies. Such antibodies which can be obtained by one skilled in art include HLe-1 which recognizes an antigen present on mature human leukocytes, such as granulocytes, 25 and very immature erythrocyte precursor but not nucleated cells, and antibodies to the transferrin receptor. Procedures using see Bianchi, supra PCT/US90/06623.) antibodies can be carried out by contacting a sample containing fetal and maternal blood with a labeled 30 antibody recognizing either fetal cells or maternal cells. The antibody labeled cell can be sorted using standard techniques including flow cytometry, immunomagnetic beads and cell panning.

Non-dividing human cells should be isolated under activation 35 mild conditions designed to prevent extracellular proteases (for those of the instance plasma), intracellular proteases, or nucleases, as well as

damage to cell prevent mechanical Inadvertent protease or nuclease activation during nuclear isolation could result in damaging both the genetic material of the cell and the protein structures within or 5 around the nucleic acid. Possible nucleic acid damage includes, nucleic acid degradation, and damage to the structural state (e.g., supercoiling). One advantage of keeping the protein structure intact, is maintaining the cytoskeletal protein so it can be subsequently separated 10 from nucleic acid under mild conditions minimizing damage to histones and non-skeletal proteins.

Preferably, solutions used to isolate cells contain protease inhibitor. Solutions used to isolate cells such as HBSS and NIB solutions can be supplemented 15 with protease inhibitors as follows: 0.1 mg/ml heparin, 0.1 mM TPCK (N-tosyl-L-phenylalanine chloromethyl ketone), 0.1 mM TLCK (Nα-p-tosyl-L-lysine chloromethyl ketone), 0.05 mM PMSF (phenylmethylsulfonyl fluoride), 5 μ g/ml leupeptin, or 31.25 mM Na₂S₂O₅.

After cell purification, the cell nucleus preferably isolated and pretreated under mild conditions. Nuclear pretreatment is preferably comprised of two steps, which may be carried out simultaneously or separately; (1) and separation or membrane permeabilization, (2) and degradation) 25 alteration (e.q., denaturation matrix proteins and nuclear cytoskeletal Treatment should be carried out to minimize the damage to nucleic acid within the nucleus.

The separation or alteration of certain protein 30 appears to be a necessary step for activation. In Xenopus erythrocytes, for instance, proteolytic digestion of cytoskeletal proteins, such as vimentin, appears to be a necessary step for subsequent nuclear activation. Coppock Developmental Biology 131:102 (1989). pretreatment should prepare the nucleus for subsequent activation rather than cause activation.

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conditions Desired for plasma membrane permeabilization include mild detergent treatment, mild protease treatment, mild shearing, and mild hypotonic Mild conditions are those conditions able to shock. permeabilize the plasma membrane while creating the least amount of damage to the nuclear DNA and proteins. Permeabilization can be detected using trypan blue. Trypan blue is a dye which cannot enter intact cells. cell indicates entrance of trypan blue into а 10 permeabilization. Protein degradation due to inadvertent protease activation can be determined using polyacrylamide gel electrophoresis to look for protein degradation products. The intactness of nuclear nucleic acids can be established by using agarose gel electrophoresis to determine the presence of nucleic acid degradation products.

separation Possible pretreatments for or alteration of cytoskeletal proteins and nuclear matrix proteins include the following:

- 20 (a) Treatment with one or more thiol reducing agent such as 10 mM dithiothreitol for a limited time, at a controlled temperature and pH, to denature cytoskeletal protein;
- Controlled salt extraction, such as 25 washing in buffers supplemented with increasing amounts of NaCl or KCl in the range of 0.025 to 1.0 M, to selectively remove cytoskeletal proteins and proteins bound to DNA;
 - (c) Controlled poly-anion treatment, such as heparin at 0.01-1.0 mg/ml or penta sodium tripolyphosphate at 70 mM, in 10 mM borate buffer (TPP) at pH 9.0, to cytoskeletal selectively remove positively charged protein;
 - (d) Degradation of cytoskeletal proteins using a protease.
- 35 The extent of protein and DNA damage can be measured as described above. Preferably, isolation and pretreatment are both accomplished at the

same time using mild concentrations of lysolecithin (e.g., 40 μ g/ml) and protease (e.g., 0.3 μ g/ml trypsin), such that a minimal amount of damage to non-cytoskeletal The minimum proteins, histones, and nucleic acid occurs. 5 time and temperature required for detergent and protease treatment should be used. In the case of red blood cells this is about 10 minutes at 25°C, using 0.3 μ g/ml of trypsin and 40 μ g/ml of lysolecithin. As would be time one skill, the preferred and appreciated by 10 temperature will change as the concentration of reagents change.

Controlled treatment with ion-selective chelating agents may also be performed as an additional pretreatment. Suitable ion-selective chelating agents include EGTA which can chelate Ca²⁺, EDTA which can chelate Ca²⁺ and Mg²⁺, and mimosine which can chelate of Cu²⁺, Al³⁺, and Fe³⁺. These ions stabilize higher order chromatin structure, thus their chelation may aid in chromatin decondensation.

Methods to terminate the detergent and protease

20 pretreatment include adding proteins to adsorb detergents
(such as 0.4% bovine serum albumin, the bovine serum
albumin employed at this step should be prepared by
dialysis of commercially available BSA fraction V against
distilled water to remove soluble salts followed by

25 lyophilization), and adding protease inhibitors (such as
soybean trypsin inhibitor) to the reaction. The
pretreated nuclei should be subsequently washed using an
ice cold solution designed to preserve genomic DNA
intactness. NIB buffer can be used for this purpose. NIB

30 is made up of 250 mM sucrose, 25 mM NaCl, 10 mM Pipes, 1.5
mM MgCl₂, 0.5 mM spermidine, and 0.15 mM spermine, pH 7.0.

The overall efficacy of mild conditions to obtain a pretreated nucleus can be determined by: a) microscopic examination of nuclei to assess whether nuclei are free of their surrounding cytoskeleton and are free standing or clumped, clumping of nuclei is a strong indication of nuclear damage since many nuclei get trapped in released

DNA; and b) the ability of nuclei to respond to activating egg extract, the use of mild conditions increases subsequent activation of individual nuclei and improves the synchrony and homogeneity with which the entire population of nuclei is activated.

(2) Preparation Of Activating Egg Extract

Activating egg extracts can be used to cause non-dividing nuclei to swell, assemble nuclear envelopes and lamina, replicate their genomes, enter mitosis, and form 10 metaphase chromosomes. Activating egg extracts contain material, such as precursors, protein(s), nuclear membrane vesicles, or mRNA required to activate non-dividing nuclei.

Non-activated eggs can be triggered en masse to 15 produce material which brings about activation, by being chemically induced to enter the cell cycle. Eggs can be induced using standard techniques such as electric shock, pricking with a needle, fertilization and the use of a calcium ionophore. (See, Gerhart, et al., J. of Cell 20 Biology 98:1247, 1984, and the procedures described below.) The induced eggs enter into the cell cycle. appears that when an egg is at the point in the cell cycle just prior to the S-phase, the egg cytoplasm is most active in supporting activation. As the egg proceeds into 25 and past the S-phase, it appears to produce material inhibitory to nuclear activation (see Table 1).

One of the benefits of the disclosed procedures is obtaining an activating egg extract having a higher DNA synthesis activation activity than activating egg extract disclosed in the prior art. The DNA synthesis activation activity can be determined by measuring the synthesis of DNA using labelled precursors.

Hardened Xenopus eggs are a good source for preparing an activating egg extract. Hardened Xenopus eggs are stable for several hours. In contrast, "soft" Xenopus eggs must be used rapidly. As soon as soft eggs

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are dejellied they tend to induce spontaneously and This is considered disadvantageous because randomly. activating egg extracts prepared from a specific time during the cell cycle, just prior to the S-phase, have a 5 higher DNA synthesis activating activity than extracts prepared from other phases of the cell cycle. Thus, it is desirable to synchronously induce a large number of eggs which are all at the same point of the cell cycle, so extracts can be prepared later from a large number of eggs all of which have elevated DNA synthesis activation activity at the same time.

Freshly ovulated Xenopus eggs can be hardened by stabilizing the eggs vitelline envelope as described by Wangh, J. Cell Science 93:1 (1989). Obtaining freshly 15 ovulated eggs from female Xenopus is facilitated by injecting hormones which cause Xenopus to ovulate. Injecting 600 units of human chorionic gonadotropin (HCG) into a Xenopus female generally brings about ovulation within 12-15 hours. Injection of pregnant mare serum 20 gonadotropin about 24 hours before HCG treatment significantly increases the yield of mature Furthermore, repeated ovulation of frogs once every 4-8 months improves the yield of eggs by increasing the synchrony of oocyte development in the ovary.

The freshly ovulated eggs within their jelly coat, are flooded with 0.3 X NKH (1 X NKH is 40 mM NaCl, 2.5 mM KCl, 7.5 mM Hepes, pH 7.4 with NaOH), for 15-20 minutes. During this time the jelly layers swell. eggs are then dejellied in 3 X NKH containing 2% cysteine, pH 7.9, by gentle swirling for about 5 minutes.

The resulting soft eggs can be "hardened" by immediately rinsing them five times in 3 X NKH containing 2 mM MgCl₂, 1 mM CaNO₃, 10 μ M ZnCl₂ and letting them stand for at least 20 minutes at room temperature. eggs are sorted to remove damaged and partially induced Calcium is required for hardening and must be present in the 3 X NKH used to wash cysteine-treated eggs.

Eggs washed in the absence of Ca²⁺ and subsequently treated with Ca²⁺, and Ca²⁺ treatment of eggs still in the jelly coat, do not result in hardened eggs. Additionally, adding Ca²⁺ before or during dejellying will not result in hardening.

Activating egg extract is preferably obtained Induction can be from hardened eggs induced en masse. carried according to procedures described by Coppock et al., Developmental Biology 131:102 (1989). The procedure 10 described by Coppock et al. as modified, in the following manner, was used to obtain "prepared activating egg extract": 5-15 ml of hardened eggs were rinsed using activation buffer (4 mM NaCl, 0.14 mM potassium gluconate, 2 mM Hepes, 2 mM MgSO₄, and 0.6 mM Ca(NO₃)₂, pH 7.8), and 15 placed in 500 ml of activation buffer; the eggs were then induced by adding calcium ionophore A23187 (10 μ M in DMSO; Sigma Chemical Co.) to a final concentration of 100 nM and incubating at room temperature; after 10 minutes calcium ionophore treated eggs were rinsed and induced for an 20 additional 15-20 minutes by incubating in 1.5 X NKH containing 2 mM MgCl2, and 0.6 mM CaCl2 (Coppock et al., supra, stops the induction at 10 minutes); the eggs were then placed on ice in a siliconized or teflon beaker and washed 3-5X in several hundred milliliters of ice cold EB 25 buffer (EB = 50 mM potassium gluconate, 250 mM sucrose, 10 mM potassium HEPES, 1.5 mM MgCl2, pH adjusted to 7.5 with potassium hydroxide); eggs were then transferred to a volumetric polyallomer centrifuge tube, mixed with Versilube F-50 oil (General Electric) at 0.2 ml oil/ml eggs, and 30 tight packed by centrifugation at 40X g for 1 minute, at 2-4°C; the overlaying oil and aqueous layers were removed and the eggs were crushed by centrifugation 15 minutes at 9,000X g, at 2-4°C; the cytoplasmic layer between the yolk pellet and the overlaying lipid layer was collected from 35 the bottom by puncturing the tube with a syringe needle; cytochalasin B was added to a final concentration of 10-50 μ g/ml and the cytoplasmic material recentrifuged for 15

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minutes at 9,000X g, at 2-4°C; the resulting second cytoplasmic supernatant was recovered and either used fresh or frozen for future use.

This procedure for "prepared activating egg 5 extract" involving an increased induction time of 15-20 minutes over that described in Coppock et al. supra, was chosen based upon the following two experiments: plasmid DNA injected into non-activated Xenopus eggs does not begin replicating until 25-30 minutes after eggs are induced, during this lag period factors required for DNA synthesis are possibly released, altered, or synthesized within the egg; and 2) extracts prepared from eggs induced for only 10 minutes synthesize additional proteins in vitro which first act to increase DNA synthesis in pretreated Xenopus erythrocyte nuclei and then act to inhibit DNA synthesis in these same nuclei.

The second experiment, "induction optimization," particularly useful in determining the induction time for obtaining activating egg extract having an elevated DNA synthesis activation activity. experimental results for induction optimization used to obtain an activating egg extract with an elevated DNA synthesis activation activity from Xenopus, is shown in The same experimental design could be used to Table 1. establish the induction time needed to obtain egg extracts having elevated DNA synthesis activation activity from species other than Xenopus.

Induction optimization was carried for Xenopus as describe below (see also Example 2, infra, for further optimization experiments). Xenopus erythrocyte nuclei were isolated and pretreated with lysolecithin and trypsin 2 (described below). as described in Example activating egg extract was prepared from hardened eggs which were induced for 10 minutes as described above. 35 Both the activating egg extract and the pretreated nuclei were kept on ice (about 4°C). The activating egg extract was supplemented with 1 mM ATP (not used in the other

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creatine µg/ml described herein), 10 examples phosphokinase, 10 mM creatine phosphate, 10 μ Ci P^{32} -dCTP and combined with pretreated nuclei (about 200 nuclei/ μ l). Individual aliquots containing activated nuclei were 5 shifted from 4°C to 25°C. Cycloheximide to a concentration of 100 μ g/ml was added to the individual aliquots at different times. The aliquots were then incubated at 25°C for a total time, including the time at 25°C before addition of cycloheximide, of 60 minutes. After 60 10 minutes, P32-dCTP incorporation into newly synthesized DNA was determined.

TABLE 1

Time CHM Added (Minutes)	Cpm Incorporated into DNA
No CHM	1,296
0	2,894
5	4,208
15	4,937
30	3,775
45	2,314

The result of induction optimization for Xenopus activating egg extract is shown in Table 1. The highest observed DNA synthesis activation activity was 15 minutes after the addition of cycloheximide. Thus, about fifteen minutes appears to be the additional time required for peak DNA synthesis activation activity (total induction time of about 25 minutes). A more precise time point for the activation peak may be readily determined by taking additional experimental time points. Elevated DNA synthesis synthesis activation activity (more DNA 30 activation activity than zero time), was seen after an The elevated DNA additional 5, 15, and 30 minutes. synthesis activation activity decreased from 15 to 30 minute time points. After the 45 minute time point the observed DNA synthesis activation activity was below that 35 of the elevated DNA synthesis activation activity. The

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decrease in DNA synthesis activation activity observed for incubation times longer than an addition 15 minutes is attributed to the synthesis of proteins inhibitory to activation.

As would be appreciated by one skilled in the art, the optimal DNA synthesis activation time will also vary as the temperature changes. As the temperature increases the optimal DNA synthesis activation time decreases, however, the temperature is preferably not raised above 24°C. As the temperature decreases the optimal DNA synthesis activation time increases, however, the temperature is preferably not lowered below 16°C.

Several proteins present in Xenopus egg extracts are involved in DNA replication. One or more of these could be a positive acting protein synthesized during the first 25-30 minutes after activation responsible for the increase in DNA synthesis activation activity. positive acting proteins include: cyclin A, RFA single strand binding protein, cdk2 kinase, and RCC1 protein. There are also several proteins whose synthesis after the first 25 minutes could be responsible for the decrease in DNA synthesis activation activity. Possible proteins which could decrease DNA synthesis activation activity include cdc2 and cyclin B. Given the evolutionary 25 conserved nature of both the positive and negative acting proteins, and their functions, it is likely that eggs from species other than Xenopus also display an optimal time just before the start of S-phase when their cytoplasm is most active in supporting DNA synthesis activation.

As seen in Table 1, incubating for 10 minutes provided less than 60% of the peak DNA synthesis activity observed compared to the optimal DNA synthesis activity of extracts prepared from Xenopus eggs. Using the techniques described herein, the time required induction 35 obtaining egg extract having an elevated DNA synthesis activation activity (more than 70% of the peak activation)

can be obtained for activating egg extract prepared from egg sources other than Xenopus.

Activating egg extracts from "hardened" eggs may be used fresh in which case they support more than one 5 cell cycle in vitro. Alternately, these extracts may be frozen and then thawed, in which case they are able to support one or more cell cycles in vitro.

The activating egg extract is preferably made 7.5-10% (v/v) in glycerol and stored frozen in liquid 10 nitrogen, by standard techniques or by an increased rapid increased rapid The freezing technique. than technique freezes the extract faster suspending in liquid nitrogen. Increased rapid freezing can be achieved by spotting extract, made 7.5-10% (v/v) 15 glycerol, as 20 μ l droplets onto a block of aluminum immersed in liquid nitrogen.

Before use, frozen activating egg extracts are thawed rapidly at room temperature, put on ice, and if desired, supplemented to enhance activation activity. One 20 possible supplement is cyclic-AMP. The addition of 0.1 mM to 10 mM cAMP to activating egg extracts increases the activation activity of the activating egg extract, as measured by subsequent DNA replication in pretreated Xenopus erythrocyte nuclei. cAMP can be broken down by inhibitor Caffeine is an phosphodiesterase. phosphodiesterase and, thus, enhances the stability of caffeine and Thus, CAMP. endogenous and added phosphodiesterase inhibitors are possible supplements to enhance activation activity of activating egg extract. Indeed, the addition of caffeine to activating egg extract was found to increase subsequent DNA replication in activated Xenopus nuclei.

Appropriate egg extracts can be obtained from Useful guidelines for sources other than Xenopus. source to make either choosing an appropriate egg 35 activating egg extract or CSF extract are provided below. These guidelines are not intended to be a list of required

characteristics, but rather a list of considerations useful for choosing an egg source.

Useful guidelines for choosing an appropriate egg source for making egg extracts include the following:

- 1. Egg/embryo with substantial stores of activating cell cycle material are preferred. Such egg/embryos can be identified as those showing a series of rapid cell cycles, <u>i.e.</u>, cell divisions approximately once every hour as compared to once every day.
- 2. Moderate egg size is preferred. Moderate egg size represents a compromise between the cytoplasmic volume per egg and yolk mass per egg. Preferably a large yield of cytoplasm per volumetric measure of eggs is obtained.
- 15 3. A species in which female animals shed a large number of eggs is preferred as a means of increasing the amount of egg extract available from an animal, while keeping the cost of caring for the animal at a minimum. However in some instance, such as the activation of 20 mammalian somatic cell nuclei prior to transplantation into their corresponding eggs, it may be desirable to prepare extracts from mammalian eggs despite their small size and relatively small number per female.
- 4. Females of the chosen species are preferably identifiable by external characteristics.
 - 5. Females preferably breed in a reasonable period of time (at least once per year), and at a reasonable cost.
- 6. Eggs are preferably shed as single cells (e.g., not in jelly mass), or easily freed of jelly layers and other major external envelopes.
 - 7. Eggs can preferably be stabilized from activating once freed of extracellular coats. (see <u>e.g.</u>, Wangh, <u>J. Cell Science</u> 93:1 (1989)).
- 8. Females preferably produce high quality eggs which are uniform and regular. These features minimize waste and help in developing automated methods to sort

good and bad eggs. Some eggs, such as those of echinoderms and mollusks, are transparent and contain a prominent germinal vesicle nucleus which can be used to judge egg quality. Other eggs, such as those of Xenopus, are not transparent, but have two distinct colors which can also be used to judge egg quality.

- 9. Eggs are preferably chemically inducible in a synchronous manner such that a number of eggs may be induced at the same time and be approximately at the same 10 point in the cycle at a specified later time (preferably at meiotic metaphase state or mitotic metaphase state). In this way, extracts may be obtained from a number of eggs at the same point in the cell cycle by inducing all the eggs at one time and using all the eggs to prepare an extract at a later time.
 - 10. Females can preferably be chemically induced to ovulate thereby making it possible to increase the production of eggs from a given female.
- 11. Female are preferably not harmed by the egg collection method. Alternately, if egg collection does harm the female those females for which a commercial use of the carcass exists are preferred.
- 25 with concomitant DNA replication. Nuclear swelling without replication can be achieved by removal of membrane vesicles required for nuclear envelope assembly, or by inhibition of DNA synthesis (e.g., using reagents such as aphidicolin, mimosine, or DMAP), or by CSF extract supplemented with a kinase inhibitor (e.g., such as DMAP or staurosporine). Nuclear swelling with replication can be achieved using activating egg extracts such as those obtained from Xenopus eggs.
- 13. It is important that eggs used to prepare
 35 CSF extracts can be arrested in either the meiotic
 metaphase state, or in the mitotic metaphase state.
 Recovery of chromosomes, rather than interphase nuclei,

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For some requires cell cycle arrest in metaphase. species, extracts in metaphase arrest can be prepared directly from non-activated eggs, such as unfertilized Xenopus eggs, or can be made to cycle into and arrest in 5 meiotic metaphase. Useful reagents for bringing about and causing arrest in meiotic metaphase include cyclin \(\text{\text{090}} \) (a non-degradable form of sea urchin cyclin), other cyclin related peptides, small amounts of CSF extract (prepared from non-activated Xenopus eggs), components found in non-10 activated Xenopus eggs (such as c-MOS kinase) or Calyculin A used on echinoderm eggs (Tosuji et. al. Proc. Natl. Acad. Sci. 89:10613 (1992)).

(3) CSF Extract Treatment Of Nuclei

Non-activated CSF extract can be used to aid subsequent nuclear activation of non-dividing mammalian cell nuclei, including human cell nuclei, without directly causing nuclear swelling or DNA replication; or to directly cause nuclear swelling as discussed in section II Nuclei in CSF extract appear to condense into infra. chromosome like structures and may become surrounded by a spindle apparatus. Nuclear activation prior to contact activating egg extract is disadvantageous. Problems with premature activation include a decrease in 25 the enhancement of activation and different nuclei being activated at different times.

The ability of CSF extract to enhance activation may be increased by various supplement. In addition, the incubation conditions of nuclei in CSF extract can be 30 adjusted to improve the ability of such extracts to enhance activation of the nuclei upon subsequent contact with activating egg extract.

The CSF extract is preferably prepared from noninduced eggs arrested at meiotic metaphase. CSF extract 35 prepared from non-induced eggs arrested at meiotic metaphase contain high levels of mitosis promoting factor (MPF) activity and cytostatic factor (CSF) activity. CSF

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and MPF are factors present in CSF extract which are believed to aid in subsequent activation of quiescent nuclei by altering cytosketal proteins, nuclear matrix proteins, and nuclear histones, particularly by phosphorylation of these proteins.

MPF is an activity controlling nuclear entry into mitosis and initiation of spindle assembly. MPF is composed of two catalytic subunits, p34^{cdc2} and cyclin B. At the onset of anaphase, cyclin B is destroyed resulting in the inactivation of MPF. During anaphase the chromosomes move towards the two opposite poles of the spindle apparatus and subsequently decondense.

CSF is an activity responsible for metaphase arrest in unfertilized vertebrate eggs. CSF activity is due to at least two kinases: mitogen-activated kinase (MAP) and cdk2/cyclin (cdk2 is a kinase related to cdc2, but the regulatory subunit of cdk2 is cyclin E (or A) rather than cyclin B). The activities of MAP appears to be controlled by additional kinases such as c-Mos kinase.

One reason for obtaining CSF extract from eggs arrested at meiotic metaphase, is that both MPF and CSF are inactivated upon initiation of the cell cycle.

CSF extracts from non-induced Xenopus eggs can be prepared by a method based on the work of Lohka and Masui, 25 <u>Developmental Biology</u> 103:434 (1984), as well as that of Murray et al., Nature 339:280 (1989). A procedure for obtaining CSF extract is as follows. Eggs are obtained from one or more ovulating frogs as described above. Each batch of freshly ovulated eggs, about 500 to 1000 eggs, is 30 hardened as described above. Damaged and activated eggs are removed. The remaining eggs are combined into a large siliconized glass or teflon beaker and washed 4-5 times at room temperature (about 21°C) in approximately 500 ml EBbuffer containing 5 mM potassium EGTA, pH 7.5, (EB = 50 mM potassium gluconate, 250 mM sucrose, 10 mM potassium HEPES, 1.5 mM MgCl2, pH adjusted to 7.5 with potassium hydroxide). The eggs are then transferred to a volumetric

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polyallomer centrifuge tube, mixed with Versilube F-50 oil (General Electric) at 0.2 ml oil/ml eggs, and are tight packed by centrifugation at 40X g for 1 minute, at room temperature. The overlaying oil and aqueous layers are 5 removed and the eggs are crushed by centrifugation for 15 minutes at 9,000X g, at 2-4°C. The cytoplasmic layer between the yolk pellet and the overlaying lipid layer is collected from the bottom by puncturing the centrifuge tube with a syringe needle. Cytochalasin B is added to a final concentration of 10-50 μ g/ml and potassium EGTA is added to a final concentration of 1 mM. The cytoplasmic material is mixed by gently pipetting or rocking back and forth, the cytoplasmic material is then centrifuged for 15 minutes at 9,000X g, at 2-4°C. An alternative cen-15 trifugation procedure involves preparation of a high speed supernatant by centrifugation at >100,000X g for 2 hrs at In either case, the resulting second cytoplasmic supernatant (hereinafter "prepared CSF extract") recovered and is either used fresh or is made 7.5-10% in glycerol and frozen for future use in the same manner as activating egg extract. Preferably, CSF extract incubated at 25° C for 2 hours prior to freezing. level of histone H1 kinase activity increases several fold during the period of incubation.

Frozen extracts can be used by thawing rapidly at room temperature and then placing on ice. Thawed extracts are preferably supplemented with an ATP regenerating system consisting of 10 mM creatine phosphate, and 10 μg/ml creatine phosphokinase.

The histone H1 kinase activity, the structural state of plasmid DNA added to the CSF extract, and the inability of CSF extract to cause nuclear activation, demonstrated that "prepared CSF extract" was arrested in meiotic metaphase. The histone H1 kinase activity of the 35 CSF extract either before or after freezing was high. Upon activation of the extract with 1.2 to 4 mM Ca2+, the histone activity decreased. Preferably, 1.2 mM Ca2+ is

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used when CSF extract is supplemented with 1 mM EGTA to achieve recycling. No recycling occurs when 3 to 4 mM of Ca2+ is used in CSF extract supplemented in the presence of Negatively supercoiled circular plasmid DNA Lysolecithin-trypsin 5 added to the extract relaxed. pretreated Xenopus erythrocyte nuclei added to CSF extract failed to swell or synthesize DNA.

After further pretreatment in CSF extract, the nuclei may be activated by adding 9 volumes of "prepared 10 activating egg extract." DNA replication, measured by incorporation of labelled nucleotides into DNA strands, may be used to determine the extent to which prior treatment in CSF extract enhances nuclear activation in Labelled nucleotides useful activating extract. measuring nuclear DNA replication include microcurie amounts of P32-dCTP for radioactive measurement of newly synthesized DNA, 16-50 μM biotinylated-dUTP or BrdUTP for fluorescent measurement of newly synthesized DNA, and 250 μM BrdUTP for density labelling of newly synthesized DNA.

Several supplements to CSF extract, in the proper concentration, increased the ability of CSF extract to enhance activation activity without resulting in premature DNA synthesis activity. Useful supplements include etaglycerol-PO4, Ca2+, and protein kinase inhibitors. addition of β -glycerol-PO $_4$ increased the rate at which negatively supercoiled DNA relaxed in CSF extract and was subsequently assembled into chromatin. A concentration of about 80 mM β -glycerol-PO $_4$ was found to aid in chromatin assembly without causing DNA synthesis. Beta-glycerol-PO₄ is an inhibitor of phosphatase activity and may act by increasing the level of the phosphorylated functionallyactive form of topoisomerase II in the CSF extract.

Similarly, the addition of 100 μM Ca^{2+} increased both the rate negatively supercoiled DNA relaxed in CSF 35 extract and rate of subsequent assembly into chromatin. Calcium is a cofactor for calcium calmodulin activated protein kinases and may also act by increasing the level of phosphorylated active topoisomerase II activity in the CSF extract. The addition of 100 μ M CaCl₂ to thawed CSF extract failed to trigger its entry into the cell cycle as judged by continued high levels of histone H1 kinase activity. The addition of 100 μ M Ca²⁺ also increased both the amount and the rate of DNA synthesis in erythrocyte nuclei after addition of activating egg extract. CSF extract responded to the addition of 1.2-4 mM Ca²⁺ by increasing the rate and extent of chromatin assembly over that seen upon addition of 100 μ M Ca²⁺. However, the higher concentration of calcium also activated the CSF extract.

The association of factors, whose presence increase DNA synthesis activity of CSF extract was also examined. Apparently, one or more factors in CSF extract which aid in subsequent DNA replication are loosely held by the nuclei in CSF extract and are lost during washing. Xenopus cell nuclei were pretreated with trypsin and lysolecithin, added to CSF extracts to a concentration of 1000-2000 nuclei per µl, and either washed by diluting into excess NIB buffer and centrifuging, or not washed. Subsequent addition of activating egg extract, to a concentration of 100-200 nuclei per µl, resulted in less DNA replication for washed nuclei. For this reason, CSF extracted treated human nuclei are preferably not washed prior to contact with activating egg extract.

Nuclear activation upon contact with activating egg extract can be increased by manipulating the conditions in which nuclei are incubated in CSF extract during further pretreatment. Useful manipulations can be obtained by regulating the incubation period and temperature. The use of a warm-then-cold regime stimulates subsequent nuclei activation. Both warm and cold steps appear to exert positive effects on subsequent nuclei activation. Preferably, the warm-then-cold regime comprises incubation at about 25°C for 30-90 minutes followed by incubation at 4°C for 30-90 minutes.

Trypsin and lysolecithin treated Xenopus red blood cell nuclei incubated in frozen/thawed CSF extract using a warm-then-cold regime and contacted with fresh extensive extract resulted in activating egg 5 synchronous nuclear envelope formation, swelling, and replication upon contact with freshly prepared activating egg While this system is attractive to aid in activation of human nuclei, because of the convenience of using frozen CSF extract, use of this system on Xenopus 10 erythrocyte nuclei revealed several limitations. limitation is the need to freshly prepare activating egg extract, which is experimentally inconvenient.

Using CaCl₂ in conjunction with frozen The use of CaCl2 permits overcomes this limitation. swelling, formation, envelope 15 synchronous nuclear replication, entry into mitosis (including formation of chromosome-like structures without DNA fragmentation), and renewed DNA synthesis in a second S-phase when both frozen CSF and frozen activating egg extracts are used. 20 the use of both a warm-then-cold regime and CaCl2 is particularly advantageous when frozen activating egg extracts and frozen CSF extracts are used to cause nucleus Preferably the CSF extract contains 0.1 to activation. 0.4 mM $CaCl_2$ to enhance nuclear activation upon subsequent 25 contact with activating egg extract. At this range of Ca2+, nuclei treated in CSF extract should not activate until contact with activating extract.

(4) Activation Of Nuclei With Activating Egg Extract

Activating egg extracts can be used to activate
non-dividing mammalian cell nuclei, such as non-dividing
human cell nuclei, to bring about swelling, chromatin
decondensation, DNA replication and formation of metaphase
chromosomes. However, nuclear activation can be stopped
at various points and information about nucleic acid
sequence and structure can be obtained by examining the
resulting DNA. Under duplication conditions, the nucleus

swells, DNA replicates, and the resultant chromosomes divide. Under non-duplication conditions, the nucleus swells, DNA is replicated, but the resultant chromosomes do not divide. Under non-synthesis conditions the nucleus swells, but DNA is not replicated, and nuclei do not divide.

The use of nocodazole, or other drugs like colchine, colcemid, and D_2O which inhibit microtubule assembly is preferred for preventing separation of mitotic chromosomes. These drugs prevent the formation of mitotic spindles during the cell cycle. As a result, condensed chromosomes accumulate rather then separate to the cells poles and are readily visualized for karyotypic analysis.

However, the addition of 5 μ g/ml nocodazole to 15 activating egg extract decreases the rate of DNA Thus, to maintain a high rate of DNA replication. replication it is necessary to either: 1) use nocodazole at a dose less than 5 $\mu \mathrm{g/ml}$, such as adding nocodazole to CSF extract at 5 $\mu \text{g/ml}$ and diluting the mixture with 9 20 volumes of activating egg extract; 2) use another drug such as colchine, colcemid or D_2O (deuterium oxide) which may be able to block mitotic spindle formation without inhibiting DNA replication; or 3) add the spindle inhibitor later, i.e., after DNA synthesis is complete but before nuclei proceed into mitosis. 25

To avoid artifacts such as chromosome fragmentation during nuclear activation it is desirable that complete, rather than partial, replication of nuclear genomes be achieved. The following techniques are useful to assess the extent of genome replication achieved during nuclear activation:

1) Coordinate observations of the kinetics of DNA synthesis, the size of the DNA molecules made, the timing of mitosis following DNA synthesis, and the morphological appearance of nuclei. Complete replication is characterized by a early onset and rapid rate of DNA synthesis in all nuclei, an abrupt cessation of DNA

synthesis in all nuclei, followed by rapid entry into mitosis, and renewed replication when nuclei exit mitosis. In addition, newly synthesized DNA molecules are very long (greater than 50,000 base pairs), but are transiently cleaved by type II topoisomerase during the period of chromosome condensation and decondensation.

- 2) The isotope dilution technique can be used to measure the pool size of DNA precursors in the activating egg extract to establish the extent of genome replication, on the basis of the radioactive specific activity of the DNA. The isotope dilution technique can be carried out according to Blow and Laskey, Cell 47:577 (1986).
- 3) BrdUTP density labelling of newly replicating DNA followed by isopycnic centrifugation in CsCl and Southern hybridization can be used to determine if one or more rounds of replication is occurring. During the initial round of semi-conservative replication, incorporation of BrUTP leads to formation of a DNA duplex having one heavy (BrUTP containing) strand and one light strand. The subsequent production of a DNA duplex containing two 20 one round heavy strands indicates more than replication.
- 4) DNA replication can be visually measured using biotinylated-deoxynucleotide triphosphates (such as 25 biotin-11-dUTP) or bromodeoxy-UTP. These nucleotides can be added to activating egg extracts and are incorporated into DNA during replication. containing the labelled DNA can be recovered and examined in a fluorescent microscope. The labelled 30 conveniently visualized by staining with Texas streptavidin (for biotin samples) or FITC (fluorescein) anti-BrdUTP antibodies. Total DNA can be visualized using a fluorescent intercalating dye (such as propidium iodide or Hoechst stain) or a fluorescently tagged reagent. some cases it may be desirable to treat nuclei in the microchamber microscope slide with high salt solutions to stretch the DNA across the glass surface before DNA

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staining. A fluorescent microscope can be employed to establish whether all regions of the nuclear DNA (stained for instance with Hoechst) contain newly synthesized DNA (stained for instance with biotin-Texas Red streptavidin).

5 II. USE OF A CSF EXTRACT TO CAUSE NUCLEAR SWELLING

Nuclear swelling can be brought about in non-dividing nuclei by using a modified CSF extract, made by high or low speed centrifugation, or by using a CSF extract or modified CSF extract made by high speed centrifugation (a "partially purified CSF extract"). Use of a CSF extract to induce nuclear swelling is preferably carried out on isolated nuclei pretreated to separate the nuclei from its surrounding cytoskeleton, preferably with detergent and a protease as described herein. Preferably, the modified CSF extract is a partially purified CSF extract and is modified by either (a) diluting with an aqueous solution and/or (b) supplementing with a protein kinase inhibitor. Fresh CSF extract or frozen thawed extract can be modified.

Dilution of CSF extract to enhance its ability to cause nuclear swelling may be carried out using various aqueous solutions such as water and physiological pH buffers. The aqueous solution is preferably buffered to about pH 6.5 to about pH 7.5. An example of an appropriate buffer is EB buffer (EB = 50 mM potassium gluconate, 250 mM sucrose, 10 mM potassium HEPES, 1.5 mM MgCl₂, pH adjusted to 7.5 with potassium hydroxide). Preferably, the aqueous solution is added in an amount to achieve 25% to 75% dilution.

The ability of CSF extract to cause nuclear swelling can also be enhanced by using a protein kinase inhibitor such as DMAP or staurosporine. DMAP and staurosporine are broad range kinase inhibitors able to inhibit the actions of both CSF and MPF. Other protein kinase inhibitor able to inhibit CSF and/or MPF can be

obtained by one skilled in the art. The chosen kinase inhibitor can preferably inhibit both CSF and MPF.

Preferably, 2.5-5 mM of DMAP is used. The use of protein kinase inhibitors should block kinase activities 5 in the extract, including histone H1 kinase and result in the treated nuclei forming envelopes but failing to The proper protein kinase initiate DNA replication. inhibitor concentration can be empirically determined by one skilled in the art by measuring the extent of swelling 10 and DNA replication in the presence of different amount of protein kinase inhibitors.

Preferably, an aqueous solution and a protein kinase inhibitor are both used to modify CSF extract. Nuclei treated with diluted CSF extract supplemented with 15 DMAP (CSF-DMAP) swell to a greater volume than nuclei treated with undiluted CSF-DMAP. Preferably, containing a protein kinase inhibitor (CSF-PKH) is diluted 25 % to 75% using an appropriate buffer and nuclei are incubated for more than 60 minutes at 25°C, and more 20 preferably around 90 minutes at 25°C prior to measuring nuclei swelling.

Diluted CSF-PKH extract can be further modified by altering the Ca^{2+} and Mg^{2+} ion concentration to further increase swelling, and affect chromatin condensation and 25 decondensation. Ca^{2+} and Mg^{2+} ion concentration can be altered by addition of these ions or by removal of these ions by using chelating agents, such as ethylene diaminetetraacetic acid (EDTA) (e.g., 5 mM), or ethylene glycolbis(B-aminoethyl ether)N,N,N'N'-Tetraacetic Acid (EGTA) (e.g., 5 mM). Altering the free Ca^{2+} and Mg^{2+} ion concentration in the diluted CSF-PKH has the effect of changing the extent of nuclear swelling and the appearance of the chromatin within the nucleus. Very low or absent Ca^{2+} and Mg^{2+} ion levels enhance nuclear swelling and 35 chromatin decompaction. Increasing Ca^{2+} to 1.2 mM prevents significant nuclear swelling and chromatin decompaction. The optimal amount of Ca^{2+} and Mg^{2+} , and chelator can be empirically determined by varying the amount of chelator and cation concentration and measuring nuclei swelling.

Chelating agents or other agents which cause chromatin decondensation, such as polyanions like heparin and TPP or thiol reducing agents like DTT, need not be added directly to the diluted CSF-PKH extract. These additional agents may be used to further swell or decondense the nuclei after treatment in diluted CSF-PKH extract.

10 III. USE OF A CSF EXTRACT SUPPLEMENTED WITH A CYCLIN

A CSF extract supplemented with a cyclin can be used to induce chromosome formation without DNA replication. The CSF extract should be supplemented with a cyclin such as cyclin-090 in an amount sufficient to achieve both nuclear envelope breakdown and nuclear chromosome formation. The cyclin is expected to act by raising the level of MPF activity in CSF extracts, and thereby, induce conversion of isolated nuclei into mitotic chromosomes.

20 CSF extract supplemented with cyclin was used to activate nuclei in suspension resulting in nuclei conversion into chromosomes. However, the activation also resulted in intermingling of the formed chromosome, and the chromosomes were stretched and sheared. The use of CSF extract supplemented with cyclin-090 may be improved by sticking trypsin treated nuclei to a glass surface prior to incubating with modified CSF.

IV. ACTIVATION OF MAMMALIAN SPERM

The present invention also features a method for activating mammalian sperm, which is particularly suitable for the activation of human sperm. The method involves the pretreatment of human sperm with a protease, then activating the sperm with an activating egg extract. The present disclosure is believed to be the first describing the use of trypsin and an activating egg extract to

activate a human sperm. The pretreated sperm can be activated using activating egg extract, or the various procedures described herein, such as using a modified CSF extract (e.g., supplemented with an aqueous solution and/or a protein kinase inhibitor, or supplemented with a cyclin) to achieve nuclear swelling or chromosome formation, and using both a CSF extract and activating extract to cause nuclear activation.

The preferred method for activating a sperm involves (1) pretreatment involving a protease, a detergent, a thiol reducing agent, and preferably a thiol blocking to prevent reassociation of sulfhydryl groups; (2) further pretreatment using CSF extract; and (3) activation using an activation extract. Sperm can be obtained using techniques known in the art. Pretreatment can be carried out using a protease and a detergent either sequentially, or at the same time, followed by a thiol reducing agent, followed by a thiol blocking agent.

An example of a preferred protocol is as follows:

1. Lyse sperm in 100 μ g/ml lysolecithin for 5 min at 25°C.

- 2. Treat with 100 μ g/ml trypsin for 5-15 minutes (10 minutes is optimum) at 25°C.
- 3. Stop the lysolecithin and trypsin treatment by using 30 μ g/ml soybean trypsin inhibitor and 0.4% bovine serum albumin.
- 4. Incubate sperm nuclei in 5 mM dithiothreitol for 60 minutes at 4°C.
- 5. Stop reaction 4 by incubating nuclei in 1 mM N-ethylmaleimide for 10 minutes at 25°C.
- 6. Incubate nuclei in CSF extract for 90 min at 25°C, followed by 60 minutes at 4°C.
- 7. Incubate nuclei in activating egg extract. The above procedure results in nuclear swelling
- 35 without nuclear envelope formation during the CSF pretreatment step (step 6), and additional swelling, nuclear envelope formation, and DNA replication during the

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activating egg extract step (step 7). Steps 1-3, or their equivalent, are all required to achieve complete swelling, nuclear envelope formation, and DNA replication.

Activation of sperm cells have various uses 5 including being used to determine whether the sperm contains a particular gene or nucleic acid sequence which can be passed on during fertilization. Such studies are useful, for example, to study the effect of aging on sperm; detect chromosomal defects; and determine whether 10 foreign genes (such as those present in the human immuno deficiency virus (HIV)) are present is sperm.

An example of the usefulness of this aspect of invention is in the field of animal breeding, particularly the breeding of transgenically modified Transgenically modified animals are usually animals. created by injecting DNA sequences into early embryos. the injected DNA integrates into the host cell genome, it may end up in the germ line of the adult animal after the animal matures. Transgenic male animals are particularly 20 desirable since they can be bred to many females. However, prior to breeding the percentage of modified germ cells, as well as the copy number and distribution of the inserted genes in each cell is not known.

The methods and reagents provided herein for activating sperm cell nuclei and examining their genetic composition, for example via in situ hybridization, make it possible to determine the percentage of sperm carrying one or more copies of the inserted gene. This information can used to access the likelihood that a particular gene will be passed on to a future generation, prior to 30 breeding the animal. Such information is desirable

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because of the time and expense required to breed an animal.

V. NUCLEAR ACTIVATION ASSAY

The procedures disclosed by the present inven-5 tion, to activate nuclei, can also be used as a general assay procedure to measure nuclear activation and the presence of a nucleic acid sequence in activated nuclei. The assay would be particularly useful to identify and purify factors present in CSF extract and to study male fertility.

A basic assay to measure DNA replication of activated cell could have the following steps: isolating a nucleus, pretreating the nucleus, further pretreating the nucleus, contacting the further pretreated nucleus containing labeled 15 with activating egg extract nucleotides, and detecting incorporation of label into Preferably, a radioactive nucleotide replicated DNA. would be used to determine activation by measuring the extent of label incorporated into newly synthesized DNA.

The assay could be tailored to aid in the purification of factors present in CSF which help prepare nuclei for subsequent activation. Specifically, the assay would be performed without the addition of CSF extracts. Rather, various fractions of CSF extract would be obtained 25 by standard purification techniques, and used instead of Those fractions which increase activation CSF extract. activity can then be further purified.

Another use of a nuclear activation assay is to study male fertility by measuring the extent of activation of human sperm under different conditions. Such studies can be used, for example, to examine techniques to preserve sperm so the sperm can be later used in in vitro fertilization, to test sperm of infertile men to identify causes of male infertility (see, Brown et al., Yale Journal Of Biology And Medicine 65:29 (1992) (not admitted to be prior art), and test possible male contraceptives.

In virtually all species of animals, sperm cells undergo two reactions, capacitation and the acrosome response, before reaching and fusing with the egg surface. After the sperm nucleus enters an egg it undergoes several changes. The nucleus swells, acquires a nuclear envelope and lamina, replicates its DNA, and eventually fuses with the female pronucleus. During this process, sperm basic proteins (histones and protamines), are exchanged for embryonic histones.

10 It appears that in order for a sperm nucleus to respond to an egg cytoplasm it must first undergo some form of proteolytic digestion. A likely site of necessary proteolytic digestion are non-histone cytoskeletal proteins. Possible contraceptives could target necessary proteolytic enzymes. The affect of the contraceptive could be determined by assaying the degree to which activation is inhibited. Possible contraceptives could also target other enzymes which may be needed for activation.

Alternatively, the assay could be used to determine conditions which result in higher levels of activation thereby finding conditions which enhance fertilization.

Specific uses of the nuclear activation assay include the following:

- Assaying sperm cell treated under different conditions of preparation, cryopreservation, capacitation and handling;
- 2) Assaying the affect of sperm cell enzymes (e.g., proteases, nucleases, phosphatases, and kinases), including the inhibition of sperm cell enzymes, on activation;
 - 3) An assay to purify enzymes affecting activation;
- 4) Assaying the sperm from infertile individuals to determine if infertility is due to problems with sperm nuclear activation;

- 5) Assaying the ability of specific drugs or reagents to enhance or inhibit activation;
- 6) Assaying the affect of inhibitors or activators of sperm cell enzymes on activation;
- 7) Assaying for the presence of a gene used to create a transgenic animal; and
- 8) Assaying for the presence of viral genome, such as HIV present in sperm.

The specific nuclear activation assay used to study fertility would be tailored to study a particular aspect of activation. For example, to assay the effect of reagents on activation the sperm should be handled and prepared under mild conditions. As discussed above mild conditions are useful in minimizing inadvertent activation To obtain sperm for the 15 of proteases or nucleases. activation assay, fresh sperm samples should be first washed in isotonic saline solution under mild conditions. The sperm can then be stored by freezing in liquid nitrogen under controlled conditions in the presence of a 20 cryoprotectant. Martin et al., in PREIMPLANTATION GENETICS, Plenum Press, New York (Verinsky and Kuliev, eds, 1991).

Fresh or frozen/thawed sperm can be treated under conditions which result in capacitation as described by Martin et al. <u>Supra</u>. The sperm membrane is then permeabilized under mild conditions as described above (e.g., lysolecithin is used to permeabilize the membrane). The nuclei are then recovered from lysed sperm by mild centrifugation in isosmotic buffer. Nuclei are preferably pretreated as described above in Section II (e.g., using a membrane permeabilizer, a protease, and a thiol reducing agent).

The nuclei can then be further pretreated (e.g., using CSF extract containing 100 μ M Ca²⁺). The further pretreated nuclei are contacted with activating egg extract and activation activity is measured using standard techniques such as using a labeled reagent (e.g., P³²-CTP)

to detect DNA replication, and microscopic visualization of nuclear swelling. Additionally, in situ hybridization can be carried out to determine the presence, number and location of particular DNA sequences. The effect of various reagents on nuclear activation can be determined by adding reagents to the sperm before or after the various individual steps of isolation, pretreatment, further pretreatment or contact with activating egg extract.

10 <u>VI. RETROVIRAL INTEGRATION ASSAY</u>

The assays described herein can be used to examine integration of proviral nucleic acid, such as from HIV, into host DNA. The assays can be carried out using a whole nucleus or a pseudonucleus. Such assays can be used to identify target sites to inhibit proviral integration, and to derive anti-viral agents directed at such target sites.

A provirus is the double-stranded DNA form of a retrovirus. It is synthesized in the cytoplasm of a cell infected with a retrovirus by reverse transcription of the viral RNA. Integration of the provirus DNA into the host cell genome is a critical step in the life cycle of all retroviruses, including HIV-1, and leads to viral expression and new virus production. Thus, by blocking viral integration, viral propagation (e.g., viral multiplication and/or viral infection) can be inhibited.

An integration assay can be performed as follows:

- 1. A cell nucleus is pretreated to separate the nucleus from its surrounding cytoskeleton to form a pretreated nucleus. The choice of cell nucleus can be varied depending on the virus studied. Preferably, the cell nucleus will be obtained from a cell which is a natural host for the virus. Examples of cells susceptible to retroviral infection include mammal and plant cells.
- 35 Preferably, a human cell nucleus is used.

- The pretreated nucleus is activated and incubated with a viral integration complex. integration complex contains the proviral double stranded DNA form of the viral RNA and material needed for viral 5 integration. Thus, the integration complex contains an integrase and may contain other viral enzymes proteins. An integration complex can be obtained by one skilled in the art using standard techniques. example, a high speed supernatant of cells infected with 10 a virus can be used as an integration complex. (Brown, et al., <u>Cell</u> <u>49</u>:347, 1987). Alternatively, an integration complex can be obtained from purified viral integrase and specific oligonucleotides having the viral sequences needed for integration (Engelman et al., Cell 67:1211, The integration complex can be added to nuclei, 15 1991). chromatin or pseudonuclei, at different points in the cell For example, incubation can take place before cycle. (e.g., prior to activation), during the time that nuclei are swelling and forming nuclear envelopes in CSF extract 20 and activating extract, or during chromatin assembly, nuclear envelope formation or replication of pseudonuclei.
- Measuring integration of the viral nucleic acid into the host nucleic acid. The measurement can be carried out using standard techniques such as through the 25 use of hybridization probes targeted to viral nucleic acid The use of hybridization probes can be facilitated by amplification techniques such as amplification. Preferably, unintegrated viral nucleic acid is separated from host nuclei acid prior to using the 30 hybridization assay probe. A separation step is useful for decreasing hybridization of probes to viral nucleic acid not incorporated into the host genome. Separation can be carried out, for example, by centrifugation of nuclei through glycerol or electrophoresis of isolated 35 nucleic acids.

Alternatively an integration assay can be carried out using a pseudonucleus. A pseudonucleus can be

constructed from a plasmid DNA template which is then used as a target for retroviral integration rather than intact activated nucleus. This approach has the advantage that the oligonucleotide size of the plasmid genome is much smaller than that of a whole eukaryotic nucleus and the sequence of the plasmid genome is either known or can be readily established.

A pseudonucleus can be constructed by adding plasmid DNA to a fresh or frozen/thawed CSF extract (e.g., at 0.1 - 20 ng/μl). This material can be used immediately or frozen for latter use. The plasmid DNA can form chromatin in the CSF extract. (For example, see Sanchez et al., Journal of Cell Science 103:907, 1992 and Wangh, Journal of Cell Science 93:1, 1989, describing such chromatin formation using plasmid FV1 dervived from type 1 BPV in intact Xenopus eggs).

The assay is carried out by activating the chromatin and measuring integration of viral nucleic acid. For example, the chromatin is diluted into an additional sample of CSF extract which is activated (e.g., by the addition of 1.2 mM Ca²⁺). Activation triggers nuclear envelope formation around the chromatin and causes the chromatin to replicate.

Thus, an activation assay can be performed using a pseudonucleus in place of a pretreated nucleus as follows: 1) forming a pseudonucleus; 2) activating the pseudonucleus, and incubating with an integration complex containing viral nucleic acid before activation or at different times after activation; and 3) measuring integration of the viral nucleic acid into the nucleic acid of the pseudonucleus.

Using an integration assay, it can be determined when viral integration occurs during the cell cycle, and if an agent is effective in inhibiting viral integration.

For example, the importance of different stages of the cell cycle in viral integration can be evaluated using CSF or activating extracts supplemented with DMAP,

aphidicolin, and inhibitors of type II topoisomerase.

DMAP and aphidicolin block DNA replication but allow nuclear swelling and chromatin decondensation to proceed. Inhibitors of type II topoisomerase block chromatin decondensation, which requires type II topoisomerase activity. Examples of the use of such drugs include the following:

- 1) If drugs such as DMAP and aphidicolin inhibit chromatin decondensation but fail to inhibit viral integration, then chromatin decondensation after mitosis is probably all that is necessary for integration. In this instance, drugs could be designed to prevent integration during or prior to chromatin decondensation.
- If integration (i.e., insertion of the 15 proviral DNA into the target genome) and circularization (i.e., insertion of the proviral DNA into itself) are both blocked by DMAP and aphidicolin, then on-going DNA synthesis is probably required for viral integration. Accordingly viral integration during DNA synthesis could 20 be targeted. If DNA synthesis is required for proviral integration, it can then be determined whether integration occurs before or after the host genome target replicated. For example, bromodeoxyuridine triphosphate (BrdUTP) can be added to reactions to increase the density 25 of newly synthesized DNA strands. Samples can then be collected at the end of the S phase when genome electrophoretically After replication is complete. removing all unintegrated viral molecules, the genomic DNA can then be cleaved with a restriction enzyme that 30 recognizes two or more sites within the viral genome. preparation can then be fractionated, for example, by CsCl density gradient centrifugation and probed for released segments of the virus. If the provirus is inserted into the host genome before replication, the viral DNA will be 35 recovered in the heavy/light density peak, along with virtually all the genomic DNA. On the other hand, if the provirus is inserted after its target sequence has

replicated, the viral DNA will be in the light/light peak. After determining the timing of integration, drugs can be designed to inhibit integration and tested.

3) If integration takes place in the presence of DMAP or aphidicolin, but not both, this would indicate that DNA synthesis, per se, is not required for integration, but cell cycle-dependent properties of the cytoplasm influences integration.

Thus, using this application as a guide, skilled in the art can identify when, during the life 10 cycle of a cell, viral integration occurs, target drugs to inhibit such viral integration, and assay whether an agent Agents which inhibits viral integration. retroviral integration may be used as therapeutic agents 15 to treat a person infected with a retrovirus (such as HIV) or prevent an uninfected person from being infected with a retrovirus. A retroviral "therapeutic agent" refers to an agent which reduces, to some extent, the in vivo propagation of a retrovirus and preferably reduces, to some extent, one or more of the symptoms associated with 20 a retroviral infection.

VII. MICROCHAMBER MICROSCOPE SLIDE

Conversion and analysis of interphase nuclei to meiotic or mitotic chromosomes is facilitated using the 25 microchamber microscope slide. Referring to Fig. 4, there is shown a microchamber microscope slide 10. The microchamber microscope slide allows very small amounts of expensive and hard to come by reagents to be used sequentially on nuclei in situ. For instance, isolated nuclei can be placed into central microchamber 20 which is formed tear-drop shaped, pretreated, swelled, converted to chromosomes, stained, and then read or analyzed without further centrifugation or complex manipulation.

A thin strip of PARAFILM wax 12, or other appropriate water resistant plastic tape or like material, is annealed to a standard microscope slide 11 or coverslip

The microscope slide 11 is generally flat and 14. rectangular-shaped with a top and bottom side. coverslip 14 is generally flat and circular-shaped with a top and bottom side. The PARAFILM strip defines three 5 wells connected by two narrow channels. Center microchamber 20 is generally teardrop-shaped with a generally rounded head end and a generally arrow shaped tail end. The volume of the microchamber is preferably between 5 μ l and 50 μ l, most ideally between 10 μ l and 20 μ l.

The head end of microchamber 20 is connected to a fill well 16 by a narrow entrance channel 18. end of microchamber 20 is connected to a drain well 24 by The volume of the resulting a narrow exit channel 22. wells is determined by the thickness of PARAFILM strip 12 15 and the size and shape of the wells; these parameters are adjustable.

Microchamber 20 is covered by a thin inverted coverslip 14. A thin inverted coverslip is best suited for use with an upright compound microscope. Other types 20 of coverslips may be used. For example, an optically thin coverslip is suited for use with an inverted compound microscope.

In the preferred mode of operation, coverslip 14 completely covers microchamber 20 leaving fill well 16 and 25 drain well 24 substantially uncovered. A sample of cells, nuclei, or other material, is pipetted to the wide part of the microchamber. The microchamber is then covered with a coverslip which is caused to adhere to the upper surface of the PARAFILM strip 12 by applying two small drops of paraffin oil. The overlaying coverslip can be siliconized to minimize sticking of water and other materials to this surface.

Microchamber 20 is filled by capillary action by placing fluid in fill well 16. Excess fluid is then 35 removed from both the fill well and drain well 24. microchamber is flushed by placing fluid in the fill well and then sucking the fluid through the microchamber by

capillary action achieved by touching blotting paper to the edge of the drain well.

The general teardrop shape enhances flushing of microchamber 20. For a 10 μ l microchamber as little as 20 5 μ l of fluid is sufficient to clear the microchamber. necessary, the coverslip 14 can be removed and the fill channel 18 and exit channel 22 sealed with a small bead of silicone stopcock grease. Material can then be recovered from the microchamber.

The microchamber microscope slide is extremely versatile. It can be sterilized, placed in tissue culture medium, and used as a growing surface for cells. Further, it is possible to increase the depth of the two side wells while leaving the microchamber shallow. Each of the side 15 wells could be covered with their own lid. One of the side wells could be filled several millimeters deep with tissue culture medium while the other well is left unfilled. Tissue culture medium would then flow through the microchamber across the cells until the two side wells 20 reach equilibrium, the exact flow rate being adjustable. Further potential uses include the following: (1) analysis analysis of isolated cells, growing cells; (2) particularly fetal blood cells; (3) analysis of cell nuclei, or other subcellular particles, organelles or 25 materials; and (4) analysis of material of non-living origin.

The microchamber microscope slidé allows analysis of material by essentially all light microscopy staining techniques including the following: (1) fluorescent 30. microscopy of incorporated precursors, antibody staining, nucleic acid hybridization techniques; (2) conventional histological staining procedures; and (3) staining based on enzymatic amplification of molecular signals.

The microchamber microscope slide also allows analysis of biological material by incorporation of radioactive precursors, followed by autoradiographic detection of the incorporated precursors.

The microchamber microscope slide also allows "on-line" microscopic observation of material being treated or altered by fluids flowing through the microchamber. In particular, the microchamber microscope slide 5 is ideally used to both isolate and analyze fetal cell from maternal blood. Isolation may be achieved by first coating the microchamber with the appropriate antibody to fetal cells, or their already isolated nuclei, and the microchamber is otherwise not sticky. The microchamber 10 itself selects and holds the fetal cells, or nuclei, while the maternal cells, or nuclei, are washed away. cells, or nuclei, can then be fluorescently tagged in situ and their positions identified even before starting in vitro nuclear swelling and chromosome formation. 15 fetal cells or nuclei, can then be activated using the appropriate treatments described herein.

VIII. ACTIVATION KITS

The technology disclosed in the present invention can be used to produce activation kits useful for clinical 20 activation of nuclei and scientific research. These kits are particularly useful for prenatal screening. an activation kit to aid in scientific research include facilitating the study of complex biochemical activities including the assembly of nucleosomes and chromatin on plasmid or viral DNA, formation of eukaryotic nuclear envelopes surrounding nuclear templates, semi-conservative replication of double stranded DNA within eukaryotic nuclei, conservative repair replication of single stranded DNA independent of nuclear envelope assembly, activation 30 of quiescent cell nuclei, nuclear envelope breakdown, condensation of chromatin into chromosomes, formation of meiotic and mitotic spindles, regulated transcription of eukaryotic genes, and protein synthesis.

A basic activation kit comprises frozen 35 activating egg extract and frozen CSF extract. These extracts are prepared based upon the methods described in

the present invention. Preferably the kit contains frozen activating egg prepared from eggs having an elevated DNA synthesis activation activity. More advanced kits contain various supplements which aid in activation. The various 5 supplements are either in separate containers present in the frozen activating egg extract or frozen CSF extract.

Preferably, these supplements are in separate containers. Useful supplements includes CaCl2, nocodazole, phosphodiesterase inhibitor (e.q., β -glycerol-PO₄, 10 caffeine), cAMP, protein kinase inhibitor (e.g., DMAP). Preferably, the activation kit contains a microchamber microscope slide.

The activation kits could also be supplemented with reagents used to study activation in general or Useful 15 determine the extent of genome replication. supplements for these activities include radioactive nucleotides, biotinylated nucleotides and different dyes (e.g., biotin-Texas Red streptavidin and Hoechst).

IX. CLONING WHOLE ANIMALS FROM SOMATIC CELL NUCLEI

present disclosed by the procedures invention, to activate nuclei, are also useful for preparing a nucleus for subsequent transplantation into an egg for the purpose of directing the development of a new Prior to nuclear transplantation, the nucleus organism. The activated 25 to be transplanted is activated in vitro. nucleus is then transplanted into an egg whose own nucleus has either been removed or functionally inactivated. egg subsequently develops into an new organism under the genetic information contained direction of 30 transplanted nucleus. Uses of cloning somatic cell nuclei include, creation of a clone of genetically identical animals, cloning animals having favorable attributes, and producing more animals which are in danger of becoming extinct.

A difficulty in cloning somatic cell nuclei from 35 mammalian species is that these nuclei are imprinted with

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patterns of gene structure and function (e.g., DNA methylation patterns) which differ from sperm and egg nuclei patterns. Thus, it is necessary to reprogram somatic cell nuclei before cloning to eliminate the different patterns. Prior activation of somatic cell nuclei in an appropriate egg extract before transplanting should allow for the necessary reprogramming to enable a transplanted nucleus to give rise to either a complete, or substantially complete new organism.

Cloning using a somatic cell nucleus comprises three steps; (1) activating the somatic cell nucleus, (2) preparing a recipient egg, and (3) transplanting the somatic cell nucleus into the egg. The first step is preferably carried out using the improved procedures, disclosed above, to activate a nucleus. Preferably isolation, pretreatment, further pretreatment, and contact with activating egg extract are preformed under conditions where the activated nucleus has a high DNA synthesis activation activity.

Preparation of a recipient egg will vary depending upon the egg source. The egg source should be treated in a manner to prevent activation before nuclear transplantation. Procedures to prepare mammalian eggs, such as those described by Martin et al. supra, are know in the art.

recipient egg Preparation of a Destruction or removal destroying the egg's pronucleus. of the egg's own nucleus guarantees that the eggs genetic material (DNA) does not contribute to the growth and 30 development of the newly cloned individual. One method of destroying the pronucleus is by using ultraviolet light as described by Gurdon, in METHODS IN CELL BIOLOGY, XENOPUS LAEVIS: PRACTICAL USES IN CELL AND MOLECULAR BIOLOGY, 36:299-309, Academic Press, California. (Kay and Peng Alternatively, the egg pronucleus can be 35 eds., 1991). surgically removed by procedures known in the art such as those described by King, in METHODS IN CELL PHYSIOLOGY

2:1-36, Academic Press, New York (D.M. Prescott, ed., 1966), and McGrath and Solter, Science 220:1300-1319 (1983).

Nuclear transplantation can be carried out by These techniques, vary depending 5 standard techniques. upon the species, and are known in the art.

It should be possible to clone Xenopus in the following manner: nuclei from Xenopus red blood cells are isolated, pretreated, and further pretreated. Nuclei are 10 then activated by contact with an activating egg extract. The nuclei are then activated to different stages in the cell cycle (e.g., S-phase, G2, etc.), and transferred to recipient prepared Xenopus eggs.

Recipient Xenopus eggs are prepared for nuclear 15 transplantation by hardening using Ca2+ (as described above), and then irradiating with ultraviolet light to destroy the egg's genome. One to two activated somatic nuclei, in 20 to 50 nanoliters are then microinjected into the Xenopus egg, into the clear cytoplasmic region that 20 lies approximately 400 microns below the animal pole of the egg. The egg is then incubated under conditions that These conditions can be permit cytoplasm rotation. conveniently obtained by floating the egg on Metrizamide. Rotation of the egg cytoplasm relative to the egg cortex the of establishment important for is dorsal/ventral axis of the developing vertebrate embryo.

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X. EXAMPLES

Example 1: Further Induction Optimization

This example describes additional experiments carried out to further determine the optimal induction time for an activating egg extract. Protein synthesis during the early part of the first cell cycle in activated eggs or egg extracts is required for preparation of an activating extract capable of efficient and complete genome replication. The required proteins can either be 10 synthesized in intact eggs before preparation of extracts or in extracts including frozen/thawed extracts. As noted above, activating egg extract should be prepared from extracts induced for more than 10 minutes to enhance DNA synthesis activation activity.

It was found that proteins synthesized during the first 28-30 minutes in intact eggs (incubated at 20°C) or during the first 60-80 minutes in a freshly prepared and activated extract (incubated at 25°C), promote subsequent DNA replication. In contrast, proteins synthesized later in the first cell cycle, i.e., after replication is The changes in DNA underway, inhibit DNA synthesis. synthesis can be detected as alterations in the time which DNA synthesis starts, the initial rate of replication, and the overall amount of replication.

The amount of CaCl₂ used to induce a freshly prepared CSF extract regulates whether or not the extract exits meiotic metaphase arrest, traverses the first interphase, and re-enters the first M-phase. As judged by measurements of histone H1 kinase activity, fresh CSF 30 extract induced by the addition of 3 mM $CaCl_2$ exits meiotic metaphase, enters interphase, but fails to enter mitosis-In contrast, CSF extract induced with 1.2 mM $CaCl_2$ exits meiotic metaphase, enters interphase, and then proceeds into mitosis-I, as indicated by a second peak in

H1 kinase activity. 35

For the purpose of comparison, extracts were prepared from eggs induced and incubated at 20°C for varying lengths of time before being crushed. cases the eggs were amassed, induced, washed, crushed, and for as described extracts were prepared activating extracts with the following modifications: (1) all tubes and pipette tips used to prepare egg extracts were first treated with 1% diethylpyrocarbonate to destroy ribonuclease activity and (2) all the steps in extract 10 preparation were carried out using plastic gloves to avoid Extracts were frozen on an ribonuclease contamination. aluminum block, chilled with liquid nitrogen. and then thawed at a later time prior to being used. DNA synthesis was followed by incorporation of P32dCTP, followed by 15 electrophoresis and phosphoimager analysis.

optimal DNA that results demonstrate The are obtained by activating extracts synthesis synchronously inducing batches of eggs and incubating them for 28-30 minutes at 20°C. Of the time periods tested the 28-30 minute extracts initiated nuclear replication 20 earliest, synthesized DNA fastest, and replicated more DNA, then egg extracts induced for 10 minutes, 22 minutes, The overall order for earlier 34 minutes or 40 minutes. nuclear replication, faster DNA synthesis, and extent of 25 DNA replication was as follows: 10 minutes < 22 minutes < 25 minutes < 28 minutes > 34 minutes > 40 minutes. Because the cell cycle of the egg is so rapid, even small differences in the length of incubation period or the temperature of incubation result in suboptimal extracts.

It was also determined that maximal replication, even in the frozen/thawed 28-30 minute extract, depends on continuing protein synthesis during the first 30 minutes of the in vitro incubation. Activating egg extract were prepared as described above, induced for 28 minutes. 35 Cycloheximide was added just prior to induction, or 30 minutes after induction. Maximal DNA replication was observed for control (no cycloheximide) and cycloheximide

added 30 minutes after induction, while zero minute cycloheximide addition resulted in significantly less DNA replication. This suggests that the proteins required for efficient replication are relatively unstable but are abundantly synthesized from mRNAs recruited onto polysomes during the first 28-30 minutes following egg induction.

Example 2: cAMP supplemented activating egg extract

The affect of cAMP on DNA replication activated Xenopus red blood cells was determined. Xenopus 10 nuclei were isolated and pretreated by a method based on Coppock et al., Developmental Biology 131:102 (1989), as follows: Xenopus blood was obtained from females by cardiac puncture and collected using a syringe half-filled with Barth's solution (88 mM NaCl, 2.3 mM KCl, 0.82 mM $MgCl_2$ and 10 mM Hepes, pH 7.4) containing heparin (10 mg/ml); the blood was immediately diluted into 10 ml of ice-cold 0.6 X SSC (1 X SSC is 0.15 M NaCl, 0.015 M Sodium citrate, pH 7.0) containing 0.1 mg/ml heparin, 0.1 mM TPCK (N-tosyl-L-phenylalanine chloromethyl ketone), 0.1 mM TLCK 20 (N α -p-Tosyl-L-lysine chloromethyl ketone), 0.05 mM PMSF (phenylmethylsulfonyl fluoride), 5 μ g/ml leupeptin, and 31.25 mM $Na_2S_2O_5$; bleeds containing clots, even small ones, were rejected; diluted blood was underlaid with 0.5 volumes of ice cold Metrizamide (refractive index of 25 1.3660 in 0.6 X SSC) and centrifuged at 180g for 10 minutes at 4°C, red blood cells pelleted below Metrizamide* while white cells banded above Metrizamide; the red cell pellet was resuspended using 0.6 X SSC and centrifuged in Metrizamide four more times to obtain erythrocytes of 30 greater than 99.9% purity; cells were washed three times in NIB and resuspended at 2 X 108 cells/ml; cells were then resuspended in NIB:glycerol (7:3) and frozen in aliquots of 100 μ l in liquid nitrogen; before using, frozen cells were thawed, diluted to 4 X 10^7 cells/ml in NIB at 23° C, 35 and added to an equal volume of NIB containing 80 $\mu g/ml$ lysolecithin (40 μ g/ml final concentration) and 0.6 μ g/ml

trypsin (0.3 μ g/ml final concentration) (the trypsin used in this example, and the other examples described herein, was Sigma brand Type XIII trypsin, TPCK treated from Bovine Pancrease, approximately 11,000 units/mg solid); after 5 minutes lysolecithin and protease treatment was stopped by adding soybean trypsin inhibitor concentration of 30 μ g/ml and bovine serum albumin to a final concentration of 0.4%; the resulting nuclei were centrifuged at 800g at 0°C for 10 minutes, washed twice in 10 NIB, resuspended with ice-cold NIB and kept on ice.

Isolated and pretreated nuclei were added at 200 nuclei/ μ l to 550 μ l thawed "prepared activating egg extract" supplemented with 5 μ g/ml nocodazole, 250 μ g/ml cycloheximide, 10 μ Ci P^{32} -dCTP, 10 mM creatine phosphate, and 10 μ g/ml creatine phosphokinase. Cyclic-AMP was then added to separate aliquots to yield final concentrations of 0.0 μ M, 0.1 μ M, 1.0 μ M, or 10 μ M.

Each aliquot was warmed to 23°C and sampled over time to determine P32-dCTP incorporation into replicated 20 DNA. At each time point, a 7 μ l aliquot was taken, frozen on dry ice, and later thawed and digested by the addition of 10 μ l replication sample buffer (80 mM Tris (pH 8.0), 8 mM EGTA, 0.13% phosphoric acid, 10% Ficoll, 5% SDS, 0.2% bromphenol blue) containing proteinase K (1.0 mg/ml) for 2 hours at room temperature. Incorporated radioactivity was analyzed by electrophoresis on a 0.8% agarose gel (50V, 20 hours) followed by vacuum drying the gel and counting on a Betascope.

As indicated by Table 2, the addition of 10 μM CAMP inhibits DNA replication in activated nuclei as 30 compared to DNA replication occurring without any cAMP. DNA replication increased with 0.1 μ M and 1.0 μ M cAMP. greater increase was seen with 1.0 μM than with 0.1 μM Thus, cAMP can be used to increase the activation 35 activity of activating egg extracts. A concentration of approximately 0.3 μM , was used in subsequent studies.

TABLE 2

5	Cpm Incorporated After X Minutes	Micromoles of Cyclic AMP Added 0.0 0.1 1.0 10				
10	0	14	14	10	43	
	45	37	51	14	34	
	90	73	68	56	31	
	135	288	290	564	35	
	180	839	1,141	2,316	85	
	240	1,556	2,945	4,484	168	
	300	2,954	2,692	5,504	571	

15 Example 3: Caffeine supplemented activating egg extract

The effect of caffeine on DNA replication in activated Xenopus red blood cells was determined. The experimental conditions used were as described in Example 1 with the following changes: the concentration of cAMP was set at 0.3 μ M and caffeine was added to the activating egg extract to a concentration of either 0.2 mM, 1.0 mM, or 5.0 mM.

As illustrated by Fig. 1, caffeine at 1.0 mM in the presence of 0.3 μ M cAMP gave the highest initial rate and extent of DNA replication in activated nuclei. Thus, caffeine can increase the activation activity of an activating egg extract.

Example 4: CSF Extract Supplemented With DMAP And Treated 30 With Activating Extract

Addition of 6-dimethylaminopurine (DMAP) to CSF extracts was used to further pretreat Xenopus erythrocyte nuclei and stimulate subsequent DNA replication in activating egg extract. DMAP can be used to inhibit nucleic acid synthesis and protein kinase activity. Xenopus erythrocyte nuclei were isolated and pretreated as described in Example 1 above, and incubated in thawed "prepared CSF extract" supplemented with 80 mM \$\beta\$-glycerol-PO4, and 5 \$\mu g/ml\$ nocodazole at a concentration of 2000 nuclei/\$\mu l\$. Further pretreatment was carried out by

incubation for 30 minutes at 4°C, then 30 minutes at 25°C, Half the samples were 4°C. minutes at supplemented with 5 mM DMAP before addition of the nuclei. After the two hours of incubation, each sample was diluted 5 with 9 volumes of activating egg extract, supplemented with 5 μ g/ml nocodazole (this dose of nocodazole slows down the rate of replication) and approximately 160 μ Ci/ml P32-dCTP. Aliquots were removed over time to measure DNA replication.

As illustrated by Fig. 2, the addition of DMAP to 10 CSF extracts enhanced the ability of the CSF extract to stimulate subsequent DNA replication in activating egg extract. DMAP decreased the lag time before the onset of replication and increased the initial rate and total 15 amount of DNA synthesis.

Example 5: Warm-Then-Cold Regime

Various warm-then-cold regimes used as part of a replication DNA further pretreatment increased Thawed Xenopus erythrocyte nuclei activated nuclei. (isolated and pretreated as in Example 1 above) were added at 2000 nuclei/ μ l to thawed "prepared CSF extract," The mixture was supplemented with 80 mM β -glycerol-PO₄. incubated using various warm-then-cold regimes. end of each incubation period samples was diluted with 9 25 volumes of "prepared activating egg extract" supplemented with 5 μ g/ml nocodazole and P^{32} -dCTP. Samples were removed over time to measure the extent of DNA replication.

As shown by the data represented in Fig. 3, the following warm-then-cold regimes stimulated subsequent DNA 30 replication: 30 minutes at 25°C, and 90 minutes at 4°C; 30 minutes at 4°C, 30 minutes at 25°C, and 60 minutes at 4°C; 60 minutes at 4°C, 30 minutes at 25°C, and 30 minutes at Incubation for 90 minutes at 4°C, and 30 minutes at 25°C was not as effective as incubation regimes that had 35 a warm period followed by a cold period. Thus, nuclei

activation is preferably performed using a warm-then-cold regime.

Example 6: Activation Using Frozen/Thawed Extracts

Activation of Xenopus red blood cell nuclei was studied using frozen/thawed CSF extracts and frozen/thawed activating egg extract. Xenopus red blood cell nuclei were isolated and pretreated as described in example 1. These nuclei were further pretreated at 2000 nuclei/μl in thawed "prepared CSF extract" supplemented with 10 mM creatine phosphate, 10 μg/ml creatine phosphokinase, 5 μg/ml nocodazole, 80 mM β-glycerol-PO₄, 100 μM CaCl₂ and incubated using a warm-then-cold format of 60 minutes at 25°C followed by 60 minutes at 4°C.

After further pretreatment, samples were diluted with 9 volumes of thawed "prepared activating egg extract" containing 10 mM creatine phosphate, 10 μg/ml creatine phosphokinase, and incubated with either: A) 200 μCi/ml P³²-dCTP; B) 16 μM biotin-11-dUTP, 16 μM MgCl₂; or C) no additions.

20 At various time intervals an aliquot of each incubation was treated as follows:

- A) P³²-labelled samples were treated with sodium dodecyl sulfate (SDS), proteinase-K, and then analyzed on agarose gels to determine DNA replication. Total incorporated radioactivity was measured using a Molecular Dynamics phosphoimager. The sizes of the radioactive molecules were observed and photographed on X-ray film.
- B) Biotin labelled nucleic acid was used to visualize replicated nuclear DNA. Biotin labelled samples were fixed by mixing into the samples approximately 40 volumes of freshly prepared 1.0 mM ethylene glycol bis-(succinic acid N-hydroxysuccinimide ester) (EGS) and incubating at 37°C for 30 minutes. Fixed nuclei were stored at 8°C for 48 hours and then centrifuged onto glass coverslips (2000 rpm, at 4°C for 15 minutes) through a 25% glycerol layer. The glycerol layer was removed and the

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samples were stained with Texas Red-Streptavidin (Gibco BRL, diluted 1:40 in PBS). Coverslips were then washed with buffered saline and stained with 1.0 μ g/ml Hoechst 33258 stain (for total DNA). Each sample was examined and 5 photographed at 60X using an Olympus optical system. Using these conditions: 1) nuclear envelopes were detected under phase optics as a dark line around the nucleus; 2) total nuclear DNA was observed under fluorescent optics as Hoechst positive (blue) staining; and 3) newly replicated 10 biotinylated DNA were detected as Texas Red positive (red) staining.

C) Samples from incubate (C) were used to measure histone H1 kinase activity during the course of the experiment.

As judged by both P^{32} -dCTP incorporation and biotinylated-dUTP incorporation, new DNA replication in erythrocyte nuclei was highly synchronous and efficient. Replication began at 30-40 minutes of incubation and was No additional completed by 80-90 minutes of incubation. 20 DNA synthesis was observed between 90-140 minutes. After 140 minutes DNA replication resumed. The initial rate of DNA synthesis in this system using a frozen/thawed activating egg extract is only slightly slower than that Furthermore, it using fresh activating egg extract. appears that replication of the entire genome was achieved.

As judged by nuclear morphology and staining, swelling was observed about 20 minutes after addition of activating egg extract (T=20). At T=20 no DNA replication DNA synthesis and biotinylated-dUTP was observed. incorporation were both first observed at T=40. swelling continued until T=80 at which time nuclear envelope breakdown and nuclear condensation Photographs of 10 or more nuclei at each time point revealed that virtually all nuclei in each sample were activated at the same time and in the same manner. Hoechst staining and biotin labelling revealed that

nuclear DNA was first highly compacted (T=0), became more diffuse during the period of swelling and replication (T=20 to T=80), and then condensed into chromosome like structures (T=100 to T=180). Nuclear envelope breakdown 5 occurred at T=100 to T=120 minutes, but mitotic spindle formation was not observed in these samples. likely due to the presence of low levels of nocodazole (0.5 μ g/ml). At T=160 many of the nuclei appeared under phase contrast to have nuclear envelopes suggesting they entered a second interphase. All nuclei at T=180 had 10 distinct chromosome-like structures indicating that they DNA synthesis entered a second mitosis. incorporation) resumed between T=140 and T=160 and then stopped at T=160 in accord with a second S-phase followed by a second mitosis. 15

Histone H1 kinase levels were low at the start of S-phase (T=40) and rose gradually thereafter. Correlations of DNA synthesis, nuclear morphology, and H1 kinase levels suggested that a threshold level of H1 kinase leading to nuclear envelope breakdown and first mitosis was reached at T=100. H1 kinase levels continued to rise until T=180, despite the fact that DNA synthesis resumed at T=140-160.

First mitosis occurred relatively early (T=100 to T=120) and was not accompanied by DNA fragmentation. These observations are consistent with the view that genome replication was complete in this experiment. Agarose gel analysis of the P32-labelled DNA demonstrated that virtually all newly synthesized DNA was initially of very high molecular weight (HMW), some of this material was then converted to pieces of a rather uniform moderate molecular weight (MMW). DNA pieces of the MMW size are not degradation products and reflect a fundamental unit of DNA packaging in condensing chromosomes. MMW may be due to experimental interruption of topoisomerase II dependent deconcatenation of replicated DNA loops.

In summary this experiment demonstrates, an in vitro system using frozen/thawed CSF extracts, and frozen/thawed activating egg extracts prepared from Xenopus eggs. In this system nuclei swell, acquire new envelopes, and cycle through at least one complete S phase followed by one complete M phase. Only a limited amount of DNA synthesis takes place in a second S-phase. This system permits highly synchronous activation and cycling of quiescent cell nuclei, and is directly applicable to the activation of non-dividing human cells such as fetal cells including keratinocytes, trophoblasts, erythrocytes and leukocytes, and sperm nuclei.

Example 7: Comparison Of High Speed Versus Low Speed CSF Extract

This example compares nuclei activation using a pretreatment in either low speed or high speed "prepared CSF extract." Activation was carried out using frozen/thawed activation extract induced for 28 minutes. Nuclei activation was assayed by measuring DNA replication and Histone H1 kinase activity.

Aenopus erythrocyte nuclei were prepared as described in Example 2 and incubated in either a low speed prepared "CSF extract" or a high speed "prepared CSF extract." In both cases nuclei were then diluted into 9 volumes of a 28 minute induced activating egg extract. For each incubate one set of samples were collected to measure histone H1 kinase activity, another set of samples containing P32-dCTP was used to measure DNA synthesis. The nuclei incubated in the high speed CSF extract replicated and progressed through the cell cycle more synchronously than those incubated in low speed CSF extract.

Example 8: Use of Diluted CSF Extract Supplemented With DMAP

This example illustrates the use of diluted CSF extract supplemented with DMAP to achieve nuclear envelope

formation and nuclear structure in the absence of DNA synthesis. Xenopus erythrocyte nuclei were isolated and pretreated using lysolecithin and trypsin as described in Example 2 above. "Prepared CSF extract" was made using 5 high speed centrifugation and frozen by spotting the extract, made 7.5-10% (v/v) glycerol, as a 20 μ l droplet onto a block of aluminum immersed in liquid nitrogen. Aliquots of the extract were thawed on ice supplemented with 10 mM creatine phosphate, 10 μ g/ml 10 creatine phosphokinase, 80 mM β -glycerol-PO $_4$, and 0.1 mM While still on ice the CSF extract received a CaCl₂. small volume (1/33rd) of DMAP to a final concentration of 5 mM, and was then diluted with different amounts of EB buffer as follows: mixture 1, 100% = extract only no EB; 15 mixture 2, 75% = 3 volumes extract + 1 volume EB; mixture 3, 50% = 1 volume extract + 1 volume EB; and mixture 4, 25% = 1 volume extract + 3 volumes EB.

Each of these mixtures was warmed to 25°C and incubated for 15 minutes. Pretreated nuclei were then 20 added in 1/10 the volume to a final concentration of 2000 Samples from each incubate were taken nuclei/ul. immediately (0), 60, 90, 120 minutes later and fixed, examined, and photographed. Nuclei treatment with diluted CSF-DMAP resulted in greater swelling than nuclei treated Nuclei treated with mixture 3 25 with undiluted CSF-DMAP. had a larger extent of nuclear swelling, nuclear envelope formation, and chromatin decondensation, than nuclei treated with the other mixtures. Additional experiments using P^{32} -dCTP and biotinylated-dUTP demonstrated that no 30 DNA synthesis took place during the process of nuclear swelling described above.

Example 9: Use of Diluted CSF Extract Supplemented With DMAP, MgCl₂ and EGTA

This example illustrates the effect of diluted CSF extract supplemented with DMAP, $MgCl_2$ and EGTA on nuclear envelope formation, swelling, and chromatin structure.

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Nuclei were treated as described in Example 7 prior to the addition of CSF extract. The CSF extract (prepared as in Example 7), while still on ice was supplemented as follows: 5 mM DMAP, 16 µM Biotinylated-dUTP and 16 µM 5 MgCl₂. The supplemented extract was diluted with an equal volume of EB buffer containing 5 mM potassium EGTA, pH 7. This mixture was warmed to 25°C and incubated for 15 minutes. Pretreated nuclei were then added in 1/10 the volume to a final concentration of 2000 nuclei/µl. Samples from each incubate were taken immediately 0, 15, 45, 60, and 90 minutes later and were fixed, examined, and photographed.

The 50% CSF-DMAP, supplemented with EGTA and MgCl₂, caused pretreated erythrocyte nuclei to rapidly swell and acquire a nuclear envelope. No biotin incorporation into DNA was observed. Thus in contrast to Example 7, swelling took place in the absence of DNA synthesis. In addition, the DNA observed in this example was more compacted than the DNA observed in example 8. The difference between this example and example 8 is likely due to the alteration of CSF extract cation concentration, and composition. For example, the EGTA may chelate the Ca²⁺ thereby lowering the Ca²⁺ while additional Mg²⁺ is added to increase the Mg²⁺ concentration.

25 Example 10: Microchamber Microscope Slide

use illustrates the example This microchamber microscope slide to analyze and activate Xenopus erythrocyte nuclei were isolated and nuclei. pretreated as described above in Example 1. These nuclei, 30 in NIB buffer, were allowed to settle onto the lower surface of several microchamber microscope slides. coverslip was placed over each sample and sealed using oil Nuclei were further pretreated using along the sides. thawed "prepared CSF extract" made 80 mM in β -glycerol-PO $_4$ 35 and supplemented with 10 mM creatine phosphate and 10 μ g/ml creatine phosphokinase. Ten microliters of CSF

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extract was allowed to flow into each well and the microchamber microscope slide was then subjected to the following warm-then-cold treatment; 30 minutes on ice, 30 minutes at 25°C, and 30 minutes on ice. After the warm-5 then-cold treatment, the CSF extract in each well was displaced by the addition of 20 μ l freshly "prepared activating egg extract" containing biotinylated-dUTP. microchamber microscope slides were then warmed to 25°C. At varying lengths of time the incubations were stopped by 10 rinsing the microchamber with 75 μ l of an appropriate buffer containing Texas red streptavidin (for detection of incorporated biotin), followed by staining with Hoechst The nuclei were stain for detection of total DNA. photographed at a magnification of 60% using fluorescent optics.

Red blood cell nuclei before pretreatment were small and compact. The majority of nuclei were separated from one another indicating they were not damaged or clumped during isolation.

Red blood cell nuclei at the end of further pretreatment in CSF extract were attached to the surface of the microchamber microscope slide and remained small and highly compact.

Red blood cell nuclei 30 minutes after addition of activating egg extract swelled dramatically, and were attached to the surface of the slide. streptavidin staining of these nuclei demonstrated the lack of DNA replication.

incubation in Nuclei after 85 minutes of activating egg extract were swollen. As seen by Texas red streptavidin staining, these nuclei were surrounded with a nuclear envelope and initiated DNA replication.

incubation fresh in 150 minutes of After activating egg extract DNA replication was complete and As a result of entering 35 the nuclei entered mitosis. mitosis, the nuclear envelopes dissembled and the DNA condensed into chromosome-like structures which remained

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attached to the surface of the microchamber microscope slide.

These results demonstrate the utility of microchamber microscope slide in nuclei activation. Using 5 the methods and products disclosed in the present activated invention nuclei were conveniently microchamber microscope slide.

Example 11: Activation Of Nuclei From Fetal red blood Cells Isolated From The Umbilical Cord

The activation of human fetal red blood cells using activating egg extract is described below. fetal red blood cells were prepared from umbilical cord blood, pretreated with lysolecithin and trypsin, contacted with activating egg extract.

Human fetal red blood cells were isolated from umbilical cord blood and fractionated into a nucleated cell fraction and a non-nucleated cell fraction described by Bianchi et al., Proc. Natl. Acad. Sci. USA 87:3279 (1990). A sample of neonatal umbilical cord blood 20 was drawn into a vacuum tube containing anticoagulants, the blood was diluted 1:1 with Hank's balanced salt solution (HBSS) (Hanks and Wallace, Proc. Exp. Biol. Med. 71:196 (1949)), layered over a Ficoll/Hypaque column (Pharmacia) and spun at 1400 rpm for 40 min at room temperature. The mononuclear cell layer was recovered and washed twice by centrifugation in HBSS. The cells were then washed several times in NIB buffer (250 mM sucrose, 25 mM NaCl, 10 mM Pipes, 1.5 mM MgCl2, 0.5 mM spermidine, and 0.15 mM spermine, pH 7.0); the resulting cell pellet 30 was suspended in NIB:Glycerol (7:3) and frozen in liquid nitrogen as 100 μ l aliquots containing 6.3 x 10⁶ cells.

Frozen cells were thawed at room temperature and put on ice, washed twice with NIB, diluted to 4×10^7 cells/ml in NIB at 23°C, and added to an equal volume of 35 NIB containing 80 μ g/ml lysolecithin and 0.6 μ g/ml trypsin; lysolecithin and trypsin treatment was halted

after 5 minutes by adding soybean trypsin inhibitor to a concentration of 30 μ g/ml and bovine serum albumin to a Isolated nuclei were added final concentration of 0.4%. directly to thawed "prepared activating egg extract" to a 5 concentration of 200 nuclei/ μ l, supplemented with 10 mM creatine phosphate, 10 μ g/ml creatine phosphokinase, 5 μ g/ml nocodazole, 0.3 mM cAMP, and 1 mM caffeine. aliquot of this sample was supplemented with P32-dCTP at approximately 200 μ Ci/ml and used to measure 10 replication. A second aliquot was sampled periodically for fluorescent microscopic examination of nuclei after fixation and staining with Hoechst dye (as described above).

Human red blood cell nuclei treated in the manner 15 described above swelled significantly during the first 90 DNA synthesis minutes and initiated DNA synthesis. continued for approximately 4.0 hours after which nuclear However, the observed kinetics of chromatin condensed. DNA synthesis indicated that complete genome replication The failure to 20 was not achieved in this experiment. achieve complete genome replication was probably due to the failure to further pretreat the isolated human nuclei in CSF extract and because activating egg extract contained a relatively high level of nocodazole, i.e., 5 $\mu g/ml.$

encountered, difficulties Despite the formation of metaphase chromosomes demonstrates that the present invention can be used to activate non-dividing The non-dividing human nuclei activated human nuclei. 30 analogously to Xenopus erythrocyte nuclei. Therefore, the various improvements described in the present invention, such as further pretreatment in CSF extract, a warm-thencold regime and the addition of 100 μM CaCl $_2$, which result in increasing the activation of non-dividing Xenopus erythrocyte are applicable to activate non-dividing human nuclei.

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Example 12: Activation of Nuclei Of Fetal Red Blood Cells Isolated From Fetal Liver

This example illustrates the use of the products and methods described herein to determine preferred 5 activation conditions and activate blood cells isolated from fetal liver. Mononucleated cells isolated from fetal liver, studied in this example, were predominately fetal blood cells as judged by their red color. The following steps were preformed:

Isolation of mononucleated human liver cells. Mononucleated human cells were isolated from human fetal liver by gently trimming the tissue and then homogenizing it between two glass slides. The cells were collected by suspension in phosphate buffered saline and 15 then transferred to a centrifuge tube. 2 ml of Ficoll was suspension which the cell layered under centrifuged at 2000 rpm for 20 minutes. The red mononuclear cells (upper layer) containing predominantly erythroid blood cells were collected, diluted with 20 phosphate buffered saline and centrifuged gently to pellet the cells. Cell pellets were resuspended and pelleted one phosphate buffered saline and time in more resuspended in RPM1 tissue culture medium.

The cells were frozen in liquid nitrogen. 25 prepare cells for freezing in liquid nitrogen, the cells were pelleted by gentle centrifugation, resuspended, and salt balanced in Hank's centrifuged again containing protease inhibitors (TPCK 0.1 mM, TLCK 0.1 mM, PMSF 0.05 mM, and leupeptin 5 μ g/ml) at 4°C. The resulting supernatant was clear indicating the absence of hemolysis. The pellet was resuspended in 1-1.5 ml of NIB containing protease inhibitors (TPCK 0.1 mM, TLCK 0.1 mM, PMSF 0.05 mM, and leupeptin 5 μ g/ml), the volume was The suspension was then spun at 1000 brought to 10 ml. 35 rpm for 10 minute at 4°C, again no hemolysis was observed, and resuspended to final volume of 5 ml in NIB containing protease inhibitors (TPCK 0.1 mM, TLCK 0.1 mM, PMSF 0.05

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The concentration of cells was mM, leupeptin 5 μ g/ml). approximately 3.55 x $10^7/ml$. The cells were then spun down, resuspended in 1.775 ml 70% NIB - 30% glycerol, and frozen as 50 μ l aliquots in liquid nitrogen.

Membrane permeabilization of nuclei. Step 2: The membrane of nuclei prepared as described in step 1 was permeabilized using lysolecithin. Frozen cells were warmed quickly, diluted with NIB, and lysed by addition of lysolecithin at a final concentration of 40 10 μ g/ml for 5 minutes at 25°C. At this point the nuclei are surrounded by a cytoskeletal matrix and do not expand or divide if contacted with activating extract.

cytoskeletal proteins of 3: Removal Step surrounding the nucleus and nuclear matrix proteins within the nucleus.

The nuclei from step 2 were treated with trypsin using variable amounts of enzyme and treatment times. Increasing the length of trypsin treatment from 0-15 minutes, at 25°C, increased the extent of DNA synthesis 20 after standard CSF pretreatment and replication in Incubation times longer than 15 activating extract. minutes resulted in decreased replication, probably due to nuclear damage and clumping. Optimal trypsin pretreatment used 0.4 μ g/ml trypsin for 15 minutes at 25°C. 25 as would be appreciated by one skilled in the art, conditions for trypsin treatment may vary depending on how the cells are washed to remove protease inhibitors added during cell preparation and the trypsin incubation These results confirm that, human cell temperature. 30 nuclei, like the Xenopus erythrocyte system, should be prepared for activation using a controlled proteolytic step.

The lysolecithin-trypsin pretreatment was stopped by adding BSA to 0.4% + soybean trypsin inhibitor to a final concentration of 30 μ g/ml followed by gentle The nuclear pellet was suspended and centrifugation. pelleted once more in 0.4% BSA and then in NIB alone.

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Step 4: Further pretreatment with CSF extract.

Washed nuclei were further pretreated in CSF extract to enhance activation. Pretreatment as described in steps 2 and 3 was not sufficient to allow swelling and 5 replication of nuclei using frozen/thawed activating This was attributed to the absence of MPF activity in frozen/thawed activating extract. pretreatment with CSF extract substantially increased responsiveness of lysolecithin/trypsin pretreated nuclei.

CSF extract further pretreatments were also preformed using varying lengths of time and temperature. Frozen/thawed CSF extract were supplemented with 10 mM creatine phosphate, 10 μ g/ml creatine phosphokinase, 80 mM β -glycerol-PO₄, and 0.1 mM CaCl₂. H1 kinase levels in such 15 extracts were high and stable for several hours.

An incubation for 60 minutes at 25°C followed by 60 minutes at 4° in CSF extract was found to give the highest amount of activation. Incubations at 25°C for longer than 60 minutes resulted in lower DNA synthesis, 20 probably because individual nuclei break up into separate chromosomes. Nuclei in these experiments were at $2000/\mu l$, but a broad range of concentrations should be equally effective.

Additional experiments carried out using Xenopus erythrocytes suggest that the 60 minute cold incubation 25 after the 60 minute warm step increases the rate of subsequent replication. Possibly, the cold disassembles spindles containing microtubules that form around nuclei during the warm step.

Thus, the CSF is preferably supplemented with 10 mM creatine phosphate, 10 μg/ml creatine phosphokinase, 80 mM β -glycerol-PO₄, and 0.1 mM Ca²⁺, and a treatment regime involving incubation for 60 minutes at 25°C followed by 60 minutes at 4°C is used in nuclei further pretreatment.

Step 5: Activation of erythroid cell nuclei. Nuclear swelling, envelope formation, and replication were carried out on human fetal liver erythroid cell nuclei

prepared using the optimal conditions described in steps The nuclei were activated by diluting the 1-4 above. further pretreated nuclei into nine volumes of prepared activating extract (prepared from eggs activated for 30 5 minutes and supplemented with 10 mM creatine phosphate, plus 10 μ g/ml creatine phosphokinase, biotinylated-dUTP or 0.2 μ Ci/ μ l α P³²-dCTP). The resulting nuclei concentration was about $200/\mu l$.

The following measurement were taken: DNA 10 synthesis was monitored using extract containing $P^{32}dCTP$ by gel electrophoresis; histone H1 kinase was measured; and samples, labelled with biotinylated-dUTP were taken for cytological analysis of DNA replication and nuclear envelope breakdown.

DNA synthesis began after a lag of 30 minutes and Little or no round-2 DNA continued until 120 minutes. synthesis occurred between 150-180 minutes, probably because nuclear envelope breakdown had not taken place. Second mitosis began between 180-210 minutes as seen by 20 the rise in H1 kinase activity, and was accompanied by fragmentation of the DNA.

Analysis of the size of the replicated DNA demonstrated that the molecules were initially very large, but at the time that DNA synthesis stopped (120 minutes) a substantial portion of the DNA was converted to a middle molecular weight (MMW) band of approximately 50 kilobases. Subsequent experiments have demonstrated that formation of this band coincides with onset of mitosis, even in the absence of a significant histone H1 kinase peak, as is the 30 case in this experiment. We believe that the MMW DNA band is an artifact generated by SDS-proteinase K disruption of the topoisomerase II-DNA complexes involved in chromosome Daughter strand deconcatenation is a condensation. mandatory part of chromosome condensation and of necessity 35 requires Type-II topoisomerase (Topo II) breakage of the replicated chromosome at many sites. Thus we believe that our in vitro conditions allow the normal G2-like period to

take place following the S-phase. Chromosome condensation and deconcatenation takes place during this period.

The process of DNA fragmentation process is distinct from chromosome condensation and MMW Fragmentation appears to occur when nuclei 5 formation. enter mitosis without having completed DNA synthesis, i.e., premature chromosome condensation.

Cytotological analysis demonstrated that little nuclear envelope breakdown occurred in this or no 10 experiment at 120-150 minutes, also in accord with the Extensive nuclear absence of a histone H1 kinase peak. swelling began about 30 minutes after incubation. Nuclear envelopes formed between 30 and 60 minutes and biotin labelling of new DNA began by 60 minutes. The intensity 15 of biotin labelling increased during the S-phase, in Maximum swelling with keeping with P32 labelling. chromatin dispersion was reached at about 90 minutes, while some condensation of chromatin took place at about 90-120 minutes although a small amount of DNA synthesis 20 was still on going. In the period 120-150 minutes there was marked chromatin condensation suggesting the onset of mitotic prophase, but nuclear envelope breakdown did not take place.

The 90 minute sample was further analyzed to determine the extent of sample homogeneity. The first 36 nuclei detected with Hoechst stain were photographed. replicated nuclei Texas red staining of biotin bleeds through into the blue Hoechst channel turning the nuclei About 40% of the nuclei failed to swell and purple. 30 failed to replicate. Almost all remaining nuclei swelled and replicated to the full extent. There were very few partially replicated nuclei. The inability of some of the nuclei to replicate was attributed to nuclear damage since the response of carefully prepared frog erythrocyte nuclei is much more homogeneous.

Experiments were also carried out to determine whether nuclear swelling was dependent on DNA synthesis,

by activating nucleic in the presence of aphidicolin (added at T=0). Biotin labelling confirmed that no replication took place in the aphidicolin treated nuclei. The results demonstrated that even in the absence of replication many nuclei swelled significantly.

Step 6: Formation of mitotic chromosomes.

Swelled and replicated nuclei were treated with Cyclin-090 or CSF extract. Cyclin-090 was added to activating extract to obtain prophase mitotic chromosomes, while CSF extract was added to obtain metaphase chromosomes and nuclear envelope breakdown.

Addition of 1/20th volume of cyclin-090 at T=100 (minutes) caused a slight but real improvement in the clarity of prophase chromosomes observed in T=120 nuclei and thereafter seemed to increase the extent of chromosome condensation. Nuclear envelope breakdown was observed at T=240 and separate, chromosome-like were released.

Addition of 1/2 volume of CSF extract at T=100 caused a rapid extensive condensation of DNA and 20 disappearance of the nuclear envelope. This state of condensation remained stable until T=240.

Example 13: Activation Of Human Sperm Nuclei Using a PPT Pretreatment

of use example illustrates the This (PPT) agent 25 permeabilization-protease-thiol reducing pretreatment to enhance activation of human sperm nuclei. Fresh semen from a healthy male donor was obtained, diluted in an equal volume of yolk test buffer (Jasjey, D.G. and Cohen, M.R. Fertility Sterility 35:205-212, 1981) and frozen in liquid nitrogen in approximately 1 ml aliquots containing approximately 1x108 sperm of which approximately 68% were motile.

On the day of the experiment, sperm samples were thawed at room temperature and washed twice in ice cold NIB buffer by centrifugation. Sperm were then permeabilized by incubation in 100 μ g/ml lysolecithin for

5 minutes at 25°C and then treated with 100 $\mu g/ml$ trypsin (e.g., a protease) for 10 minutes at the same temperature. The lysolecithin/trypsin treatment was stopped by the addition of soybean trypsin inhibitor to 30 μ g/ml and 5 dialyzed/lyophilized bovine serum albumin to 0.4% and then washed by centrifugation. The sperm were then incubated in a solution of 5 mM dithiothreitol (DTT) (e.g., a thiol reducing agent) in 5 mM in NIB for 60 minutes on ice and then post-treated with 1 mM N-ethylmaleimide in NIB for 10 10 minutes at 25°C. A final wash was carried out in NIB and the nuclei were resuspended at $40,000/\mu$ l. Two additional aliquots were prepared as described above except that in one case the DTT was omitted from the 60 minute incubation in NIB, and in the other case the trypsin was omitted during the 10 minute incubation following lysolecithin treatment.

Each of the above three samples were then added at a final concentration of 4000/μl to a frozen/thawed preparation of high speed "prepared CSF extract" that had been supplemented with 10 mM creatine phosphate, 10 μg/ml creatine phosphokinase, 80 mM β-glycerophosphate, and 0.1 mM CaCl₂. Nuclei were incubated for 90 minutes at 25°C and then for 60 minutes at 4°C.

Each of the three CSF extract treated samples

were then diluted with 9 volumes of a frozen/thawed 25

minute activated egg extract supplemented with 10 mM

creatine phosphate, and 10 μg/ml creatine phosphokinase,

plus 16 μM biotinylated-dUTP and 16 μM MgCl₂, or plus 0.2

μCi/μl αP³²-dCTP. Each sample was incubated at 25°C and

sampled periodically by either fixed staining for biotin incorporation into DNA and photographed, or fractionated on agarose gel and counted for incorporation of label nucleotides into DNA.

Cytology demonstrated that each of the three treatment regimes resulted in swelling of the sperm nuclear DNA, but only the combined lysolecithin-trypsin-DTT pretreatment procedure resulted in new nuclear

envelope formation, extensive spherical swelling of the DNA synthesis as nucleus, and new DNA synthesis. determined by P32-dCTP incorporation also demonstrated that only the combined lysolecithin-trypsin-DTT pretreatment 5 procedure resulted in significant replication. combined lysolecithin-trypsin-DTT pretreatment procedure brought more than a 5 fold increase in DNA synthesis than lysolecithin and trypsin pretreatments with lysolecithin and DTT.

Additional cytological analysis and in situ hybridization was carried out using two nucleic acid probes; one to a reiterated sequence on the X-chromosome and one to a single copy sequence on chromosome 18. lysolecithin-trypsin-DTT pretreated and activated sperm nuclei were recovered after 120 minutes of incubation in activating extract in the presence of biotinylated-dUTP. Two nuclei were stained for total DNA (Hoechst=blue), the X-chromosome (using a digoxygenin-labelled probe-green), (Texas-red biotinylated DNA synthesized newly 20 streptavidin=red), and chromosome 18 (using a biotinylated probe + Texas-red streptavidin=red dots). The intensity of the Hoechst staining and the biotinylated-dUTP staining in the first nucleus was less than that in the second This demonstrates that first nucleus had only nucleus. 25 begun in vitro replication while the second nucleus had In addition, the first replicated more completely. nucleus contained only one copy of the X-chromosome, indicating that the probed region of the chromosome had not yet replicated, while the second nucleus contained two indicating that 30 copies of the X-chromosome, In addition, two copies of replicated by this time. in the first nucleus, chromosome 18 were detected suggesting that this probe detects an early replicating sequence. Detection of chromosome 18 was obscured in the 35 in the second nucleus by the higher level of incorporated biotinylated-dUTP.

These results are consistent with the notion that both thiol reduction and protease induced changes in the sperm cytoskeleton and protamines are required for displacement of protamines and their replacement by chromatin forming histones in the egg extracts. Chromatin assembly is important in the formation of the surrounding nuclear envelope and completion of the envelope is important for initiation of DNA synthesis. These results also demonstrated that human sperm cell nuclei which have been activated and replicated in vitro can be used for genetic analysis.

Additional experiments demonstrated that the optimized protocol described above also resulted in formation of mitotic chromosomes from human sperm nuclei.

Other embodiments are within the following claims.

Claims

- 1. A method for activating a nucleus from a human fetal cell comprising the steps of:
- a) separating said nucleus from its surrounding5 cytoskeleton to form a pretreated nucleus, and
 - b) contacting said pretreated nucleus with an activating egg extract to activate said pretreated nucleus.
- The method of claim 1, wherein said fetal
 cell is selected from the group consisting of keratinocyte, trophoblast, erythrocyte and leukocyte.
 - 3. The method of claim 2, wherein said leukocyte is selected from the group consisting of neutrophil, basophile, eosinophil, and granulocyte.
- 4. The method of claim 1, wherein said separating is carried out using a protease and a non-ionic detergent.
 - 5. The method of claim 4, wherein said protease is trypsin and said detergent is lysolecithin.
- 20 6. The method of claim 1, further comprising contacting said pretreated nucleus with CSF extract.
 - 7. The method of claim 6, wherein said nucleus is activated under conditions were the nucleus and its chromosomes do not divide.
- 25 8. The method of claim 7, wherein said conditions comprise adding nocodazole to said activating egg extract or said CSF extract.
 - 9. The method of claim 8, wherein said nocodazole is in amount less than 5 μ g/ml.

- 10. The method of claim 1, wherein said nucleus is activated under conditions not suitable for nucleic acid synthesis.
- 11. A method of causing a non-dividing human nucleus to activate, using activating egg extract and CSF extract prepared from hardened eggs comprising:
 - a) incubating said non-dividing human nucleus with said CSF extract prepared from hardened eggs to form a pretreated nucleus, and
- b) contacting said pretreated nucleus with said activating egg extract wherein said activating egg extract is prepared from synchronously activated hardened eggs.
 - 12. The method of claim 11, wherein said CSF extract is frozen and thawed before use.
- 13. The method of claim 11, wherein said activating egg extract is frozen and thawed before use.
- 14. The method of claim 11, wherein said incubating is performed using a warm-then-cold regime comprising incubating at about 25°C for at least 30 minutes followed by incubation at about 4°C for at least 30 minutes.
 - 15. The method of claim 11, wherein said incubating is performed using a warm regime at about 25°C for at least 30 minutes.
- 25 16. The method of claim 11, wherein Ca^{2+} is provided in said incubating step.
 - 17. The method of claim 16, wherein said Ca^{2+} is provided in an amount greater than 100 $\mu \rm M$.

- extract, comprising the steps of hardening a plurality of eggs, simultaneously inducing said eggs; and preparing an activating egg extract from said eggs wherein said eggs are induced for a length of time such that they have at least 70% of maximal activation DNA synthesis activity.
 - 19. The method of claim 18, wherein said activating egg extract is prepared from a eukaryotic cell.
- 20. The method of claim 19 wherein said 10 eukaryotic cell is an amphibian, yeast, human, echinoderm, mollusc, fish, or chicken cell.
 - 21. The method of claim 20, wherein said eukaryotic cell is a Xenopus cell.
- 22. The method of claim 21, wherein said eggs are obtained from Xenopus and said length of time is greater than 10 minutes.
 - 23. The method of claim 22, wherein said length of time is between 25 and 30 minutes.
- 24. A method of causing a non-dividing cell 20 nucleus to swell comprising the steps of:
 - a) separating said nucleus from its surrounding cytoskeleton to form a pretreated nucleus; and
- b) contacting said pretreated nucleus with a CSF extract supplemented with an aqueous solution or a protein
 25 kinase inhibitor.
 - 25. The method of claim 24, wherein said nucleus is a human nucleus.

- 26. The method of claim 25, wherein said CSF extract is a partially purified CSF extract supplemented with an aqueous solution and a protein kinase inhibitor.
- 27. The method of claim 26, wherein said aqueous 5 solution is an appropriate buffer.
 - 28. The method of claim 27, wherein said protein kinase inhibitor is either 6-dimethylaminopurine or staurosporine.
- 29. The method of claim 28, wherein said 10 appropriate buffer is provided in an amount to dilute said CSF extract by 25% to 75%.
 - 30. The method of claim 27, further comprising the step of adding an agent to further swell or decondense said nuclei after said step (b).
- 31. A method of causing chromosome formation in a non-dividing cell nucleus comprising the steps of:
 - a) separating said nucleus from its surrounding cytoskeleton to form a pretreated nucleus; and
- b) contacting said pretreated nucleus with a CSF 20 extract supplemented with a cyclin.
 - 32. The method of claim 31, wherein said cyclin is cyclin-090.
 - 33. A method for activating or studying a mammalian sperm cell nucleus comprising the steps of:
- a) pretreating said sperm cell nucleus to form a pretreated sperm nucleus wherein said pretreating comprises (i) separating said nucleus from its surrounding cytoskeleton by permeabilizing said cell nuclear membrane and incubating in the presence of a protease and

- (ii) incubating in the presence of a thiol reducing agent; and
 - b) activating said pretreated sperm cell.
- 34. The method of claim 33 wherein said sperm 5 cell nucleus is a human sperm cell nucleus.
 - 35. The method of claim 34, wherein said activating is carried out by contact with a CSF extract, wherein said CSF extract is supplemented to induce nuclear swelling.
- 36. The method of claim 34, wherein said activating is carried out by contact with a CSF extract, wherein said CSF extract is supplemented to induce chromosome formation.
- 37. The method of claim 34, wherein said activating is carried out using an activating extract.
 - 38. The method of claim 33, wherein said sperm cell nucleus is from a transgenic animal.
 - 39. The method of clam 34, wherein said human sperm cell nucleus is obtained from either a person infected with a virus or a person not infected with said virus.
 - 40. The method of claim 39 wherein, said virus is HIV.
- 41. The method of claim 34, further comprises 25 the step of analyzing said nucleus by in situ hybridization.
 - 42. The method of claim 34, wherein said step (a)(i) is carried out using a detergent and trypsin.

- 43. The method of claim 34, wherein said pretreating step further comprises a thiol blocking agent.
- 44. The method of claim 34, wherein a CSF extract further pretreatment step is carried out between said step (a) and said step (b), comprising incubating the pretreated nuclei in CSF extract.
 - 45. An activation assay for studying male fertility comprising:
- a) pretreating a sperm nucleus to separate 10 cytoskeletal protein from nucleic acid,
 - b) activating said sperm nucleus,
 - c) measuring activation of said nucleus activated in step (b).
- 46. The activation assay of claim 45, wherein said pretreating comprises using a detergent, a protease, and a thiol reducing agent.
 - 47. The activation assay of claim 46, wherein said sperm nucleus is a human sperm nucleus.
- 48. The activation assay of claim 47, wherein 20 said activating is carried out using a CSF extract supplemented with an aqueous solution, a protein kinase inhibitor, or a cyclin.
 - 49. The method of claim 47, wherein said activating is carried out using an activating egg extract.
- 25 50. The method of claim 49, further comprising a CSF further pretreatment prior to said activating.
 - 51. A viral integration assay comprising the steps of:

- a) pretreating a cell nucleus to separate the nucleus from its surrounding cytoskeleton to form a pretreated nucleus,
- b) activating said nucleus and incubating with 5 a viral integration complex containing viral nucleic acid, wherein said integration complex is added before or after said incubating; and
 - c) measuring integration of viral nucleic acid into nucleic acid of said cell nucleus.
 - 52. The assay of claim 51, wherein said viral nucleic acid is from HIV.
- 53. The assay of claim 52, further comprising a step, between said step b) and said step c), of adding an agent which inhibits integration of said HIV.
 - 54. A viral integration assay comprising the steps of:
 - a) constructing a pseudonucleus from a defined DNA template,
- b) activating said nucleus and incubating with a viral integration complex containing viral nucleic acid, wherein said integration complex is added before or after said incubating; and
- c) measuring integration of viral nucleic acid into nucleic acid of said cell nucleus.
- 55. A product for preparing a non-dividing nucleus to activate upon subsequent treatment with activating egg extract, comprising a CSF extract prepared using an eukaryotic cell, wherein said CSF extract is supplemented with Ca²⁺.
 - 56. The product according to claim 55, wherein said Ca^{2+} is provided in an amount greater than 100 μM .

- 57. A product for causing nuclear swelling comprising CSF extract supplemented with a protein kinase inhibitor and/or an aqueous solution.
- 58. The product of claim 57, wherein said CSF extract is a partially purified extract.
 - 59. The product of claim 58, wherein said aqueous solution is an appropriate buffer.
- 60. The product according to claim 59, wherein said supplement is said protein kinase inhibitor and said 10 appropriate buffer.
 - 61. The product according to claim 60, wherein said protease inhibitor is either 6-dimethylaminopurine or staurosporine.
- 62. A product for causing chromosome formation 15 in a cell nucleus comprising a CSF extract supplemented with a cyclin.
 - 63. The product according to claim 62, wherein said cyclin is cyclin- \triangle 90.
- 64. A product for activating a non-dividing 20 nucleus comprising an activating egg extract having at least 70% optimal activation activity.
 - 65. The product of claim 64, wherein said activating egg extract is prepared from one or more eukaryotic egg.
- 25 66. The product of claim 64, wherein said activating egg extract is prepared from a plurality of Xenopus eggs synchronously induced for more than 10 minutes.

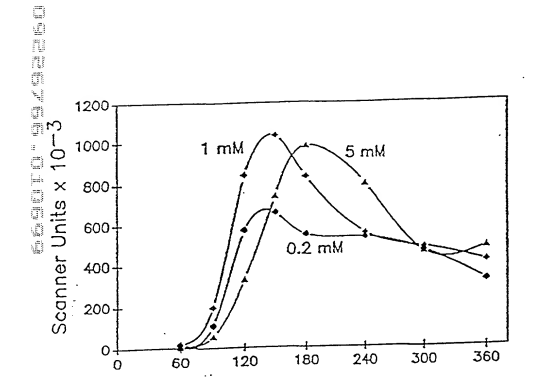
- 67. The product of claim 66, wherein said Xenopus eggs are synchronously induced for 25 to 30 minutes.
- 68. The product of claim 66, wherein said 5 activating egg extract is supplemented with cAMP.
 - 69. The product of claim 66, wherein said activating egg extract is supplemented with a phosphodiesterase inhibitor.
- 70. A kit for activating a non-dividing nucleus comprising a first product comprising frozen activating egg extract having at least 70% optimal activation activity and a second product comprising a frozen CSF extract.
- 71. The kit according to claim 70, wherein said second product is supplemented with Ca^{2+} .
 - 72. The kit according to claim 70, wherein said first product and said second product are prepared from hardened eggs.
- 73. The kit according to claims 70, further 20 comprising a microchamber microscope slide.
 - 74. A microscope slide comprising;
 - a) an upper surface,
- b) a water repellent means having a defined thickness located upon said upper surface to define a
 microchamber connected by a channel to at least one well on said upper surface, and
 - c) said microchamber shaped to enhance flushing of said microchamber.

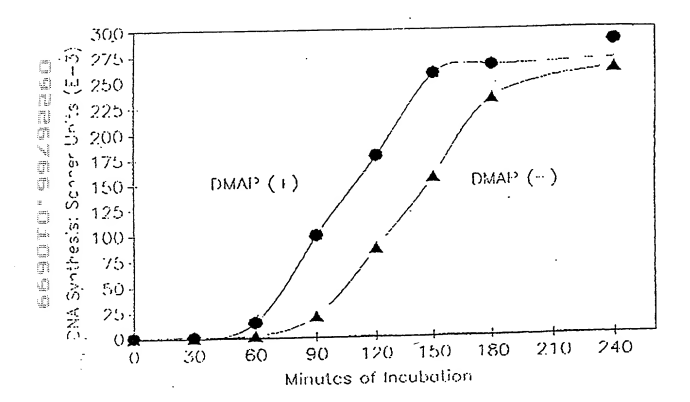
- 75. The microscope slide of claim 74, wherein said microchamber is in a teardrop-shape or a pear-shape.
- 76. The microscope slide of claim 74, wherein two wells are provided at opposite ends of said microchamber and each of said wells are connected to said microchamber by a channel.
 - 77. The microscope slide of claim 74, wherein said microchamber has a defined volume between 5 and 50 μ l.
- 78. The microscope slide of claim 74, wherein said microchamber has defined volume between 10 and 20 μ l.
 - 79. The microscope slide of claim 74, wherein said water repellent means is a tape or a coating on said upper surface of said slide.
- 15 80. The microscope slide of claim 74, wherein said water repellent means is a TEFLON® coating.
 - 81. The microscope slide of claim 74, wherein said water repellent means is a plastic containing tape.
- 82. The microscope slide of claim 74, wherein 20 said upper surface is treated to enhance cell growth compared to an untreated slide.
 - 83. The microscope slide of claim 74, wherein said microscope slide is sterile.
- 84. The microscope slide of claim 74, wherein 25 said microchamber or said at least one well contains an antibody to a human fetal cell.

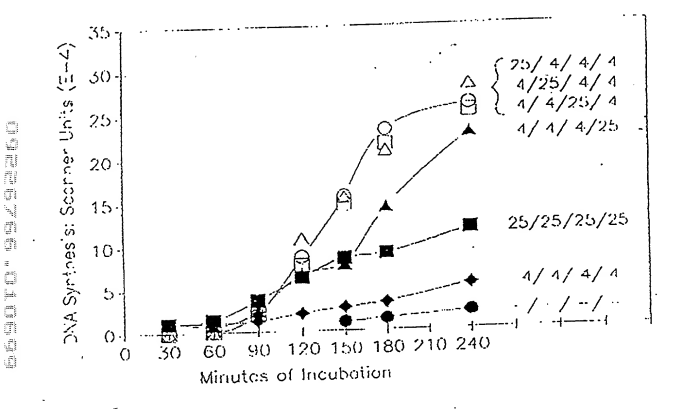
- 85. A method for analysis of growth or manipulation of a cell or cell component comprising providing a microscope slide of claim 74 and placing said cell or said cell component in said microchamber.
- 5 86. The method of claims 74, wherein a fluid is introduced into one said well and is allowed to enter said microchamber and then is removed from said microchamber.

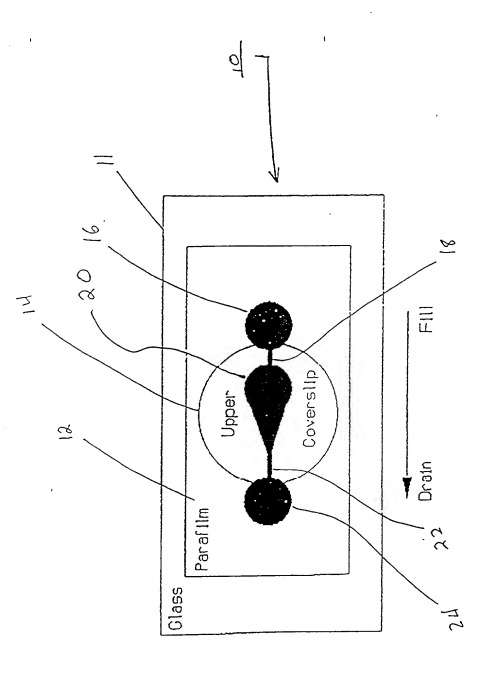
Abstract

The present invention concerns products and methods particularly useful for activating and analyzing non-dividing cell nuclei. The featured products include activating egg extracts, cytostatic factor (CSF) extracts, kits containing these extracts, and a microchamber microscope slide useful in analyzing nucleus activation.









F.g. 4

NEY

(Continuation or CIP Application,

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name,

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the

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